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* * * * * * * STN Columbus * * * * * * * * * * * * * *

FILE 'HOME' ENTERED AT 14:58:43 ON 27 AUG 2008

=> file reg

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Uploading C:\Program Files\Stnexp\Queries\10572914.str

chain nodes :
11 12 13 14 15 16 19 20 22 23 28

ring nodes : 1 2 3 4 5 6 7 8 9 10

ring/chain nodes :

17 21

Page 1

chain bonds :
1-28 2-20 7-14 8-11 11-12 11-13 12-15 12-16 14-17 14-19 20-21 22-23
ring bonds :
1-2 1-6 2-3 3-4 4-5 4-7 5-6 5-10 7-8 8-9 9-10
exact/norm bonds :
1-28 2-20 7-14 11-12 11-13 14-17 14-19 20-21 22-23
exact bonds :
8-11 12-15 12-16
normalized bonds :
1-2 1-6 2-3 3-4 4-5 4-7 5-6 5-10 7-8 8-9 9-10
isolated ring systems :
containing 1 :

G1:H,Ak

G2:SO2,[*1-*2]

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 19:CLASS 20:CLASS 21:CLASS 22:CLASS 23:CLASS 28:CLASS

L1 STRUCTURE UPLOADED

=> d 11

L1 HAS NO ANSWERS L1 STR

$$\begin{array}{c} G1 \\ N \\ O \\ H \\ \end{array}$$

1 2

G1 H, Ak G2 SO2, [@1-@2]

Structure attributes must be viewed using STN Express query preparation.

=> s 11 full

T.3 208 SEA SSS FUL L1

=> file ca

=> s 13

L42 L3

=> d ibib abs fhitstr 102

2 ANSWERS ARE AVAILABLE. SPECIFIED ANSWER NUMBER EXCEEDS ANSWER SET SIZE The answer numbers requested are not in the answer set. ENTER ANSWER NUMBER OR RANGE (1):1-2

ANSWER 1 OF 2 CA COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:373698 CA

TITLE: Preparation of 4-aminoquinoline-3-carboxamide

derivatives as PDE4 inhibitors

INVENTOR(S): Edlin, Christopher D.; Eldred, Colin David; Keeling, Steven Philip; Lunniss, Christopher James; Redfern,

Tracy Jane; Redgrave, Alison Judith; Woodrow, Michael

PATENT ASSIGNEE(S): Glaxo Group Limited, UK PCT Int. Appl., 63 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	TENT	NO.			KIN	D	DATE								D	ATE	
WO	2005	0302	 12		A1	_	2005	0407	•		004-				2	0040	923
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		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NA,	NΙ,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KΕ,	LS,	MW,	ΜZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
		ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	ΙΤ,	LU,	MC,	NL,	PL,	PT,	RO,	SE,
		SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,
		SN,	TD,	ΤG													
EP	1673	086			A1		2006	0628		EP 2	004-	7656	56		2	0040	923
EP	1673	086			В1		2008	0123									
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,	CY,	TR,	BG,	CZ,	EE,	HU,	PL,	SK,	HR	
JP	2007	5067	03		Τ		2007	0322		JP 2	006-	5273	74		2	0040	923
AT	3845							0215									
_	2298							0516									
US	2008	0096	884		A1		2008	0424									
RIORIT	Y APP	LN.	INFO	.:									2		A 2		
													844		W 2	0040	923
THER SO	OURCE	(S):			CASI	REAC	T 14	2 : 373	3698	; MA	RPAT	142	:373	698			

GI

The title compds. I [R1 = (un)substituted aryl, heteroaryl, heterocyclyl, etc.; R2 = H, alkyl; R3 = H, alkyl, cycloalkyl, etc.; R4 = H, alkyl; or NR3R4 = (un)substituted heterocyclyl; R5 = H, alkyl; R6 = H, alkyl, alkoxy, etc.] which are inhibitors of phosphodiesterase type IV (PDE4) and are of use in the treatment of inflammatory and/or allergic diseases, were prepared Thus, reacting 4-chloro-6-(1-piperidinylsulfonyl)-3-quinolinecarboxamide (preparation given) with 3-methoxyaniline afforded II. The exemplified compds. I inhibit the catalytic activity at PDE4B (human recombinant) enzyme with pIC50's in the range 7.5-10.8. The pharmaceutical compns. comprising the compound I are disclosed.

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of 4-aminoquinoline-3-carboxamides as PDE4 inhibitors) RN $849591-19-1\ {\rm CA}$

CN 3-Quinolinecarboxamide, 4-[(3-methoxyphenyl)amino]-6-(1-piperidinylsulfonyl)- (CA INDEX NAME)

Ι

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 2 OF 2 CA COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:355178 CA

TITLE: Preparation of aminocarbonylquinoline derivatives as

phosphodiesterase type IV (PDE4) inhibitors

INVENTOR(S): Edlin, Christopher; Eldred, Colin David; Lunniss,

Christopher James; Redgrave, Alison Judith; Robinson,

John Edward; Woodrow, Michael

PATENT ASSIGNEE(S): Glaxo Group Limited, UK SOURCE: PCT Int. Appl., 69 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT	NO.			KIN	D	DATE			APPI	LICAT	-			D	ATE	
WO	2005	0307	 25		A1	_	 2005	0407	1	WO 2					2	0040	927
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	, MK,	MN,	MW,	MX,	MZ,	NA,	NΙ,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	, SC,	SD,	SE,	SG,	SK,	SL,	SY,
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	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	, SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
		AZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
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		SN,	TD,	TG													
EP	1673	345			A1		2006	0628		EP 2	2004-	7686	49		2	0040	927
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		IE,	SI,	LT,	LV,	FΙ,	RO,	CY,	TR,	BG,	, CZ,	EE,	HU,	PL,	SK,	HR	
JP	2007	5067	17		T		2007	0322		JP 2	2006-	5274	83	•	2	0040	927
US	2007	0191	426		A1		2007	0816	1	US 2	2007-	5729	13		2	0070	206
PRIORIT	Y APP	LN.	INFO	. :					(GB 2	2003-2	2272	6		A 2	0030	927
									1	WO 2	2004-0	GB41	06		W 2	0040	927
OTHER S GI	OURCE	(S):			CASI	REAC	T 14	2:35	5178	; M.	ARPAT	142	:355	178			

AB Title compds. I [R1 = (un)substituted-aryl, -heteroaryl, cycloalkyl, etc.; R2 = H, alkyl; R3 = H, (un)substituted alkyl, cycloalkyl, etc.; R4 = H, alkyl; or R3 and R4 together = (un)substituted N-heterocycle; R5 = H, alkyl; R6 = H, alkoxy, Cl, etc.] and their pharmaceutically acceptable salts, are prepared and disclosed as inhibitors of phosphodiesterase type IV (PDE4). Thus, e.g., II was prepared by amidation of 3-(aminocarbonyl)-4-{[3-(methyloxy)phenyl]amino}-6-quinolinecarboxylic acid (preparation given) with morpholine. The inhibition capability of I was evaluated in radioactive Scintillation Proximity Assay (SPA) and revealed that selected compds. of the invention possessed pIC50 values in the range of 6.3-9.5. I as PDE4 inhibitors should prove useful in the treatment of inflammatory and allergic diseases.

IT 849124-91-0P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of aminocarbonylquinoline derivs. as phosphodiesterase type IV (PDE4) inhibitors)

RN 849124-91-0 CA

CN 3-Quinolinecarboxamide, 4-[(3-methoxyphenyl)amino]-6-(4morpholinylcarbonyl)- (CA INDEX NAME)

Ι

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> file marpat

=> s 11 full

FULL SEARCH INITIATED 14:59:50 FILE 'MARPAT'

FULL SCREEN SEARCH COMPLETED - 9902 TO ITERATE

100.0% PROCESSED 9902 ITERATIONS

SEARCH TIME: 00.00.07

L5 131 SEA SSS FUL L1

=> d ibib abs fqhit 1-75

L5 ANSWER 1 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 149:128754 MARPAT

TITLE: Preparation of 8-hydroxyquinolines for treatment of

neurological conditions

INVENTOR(S): Barnham, Kevin Jeffrey; Gautier, Elisabeth Colette

Louise; Kok, Gaik Beng; Krippner, Guy

131 ANSWERS

PATENT ASSIGNEE(S): Australia

SOURCE: U.S. Pat. Appl. Publ., 120pp., Cont.-in-part of U.S.

Ser. No. 521,902.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATE	I TN	. O <i>l</i>		KII	ND	DATE			A	PPLI	CATI	ON NO	Э.	DATE			
									_								
US 2	0080	0161	353	A.	1	2008	0703		U	S 20	07-9	0194	1	2007	0919		
WO 2	0040	0074	61	A.	1	2004	0122		W	0 20	03-A	U914		2003	0716		
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	RW:	GH,	GM,	ΚE,	LS,	MW,	ΜZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,

KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG US 20060089380 US 2005-521902 Α1 20060427 20050810 IN 2006-KO1346 IN 2006KO01346 Α 20070720 20061211 PRIORITY APPLN. INFO.: AU 2002-950217 20020716 WO 2003-AU914 20030716 US 2005-521902 20050810 IN 2005-KN166 20050210 GΙ

Ι

AB The title compds. with general formula I [wherein R2 = (un)substituted alkyl, alkenyl, aryl, heterocyclyl, etc.; R, R1, and R3 = independently H, OH, cyano, (un)substituted alkyl, etc., with the proviso that when R and R1 are H and R2 is COOH or CO-OMe, then R3 is not OH.] or pharmaceutically acceptable salts, hydrates, or solvates thereof were prepared for the treatment of a neurol. conditions. For example, 5,7-dichloro-8-hydroxyquinoline-2-carboxylic acid (preparation given), dicyclohexylcarbodiimide, 1-hydroxybenzotriazole hydrate, histamine dihydrochloride, and Et3N were stirred in DMF/CH2Cl2 to give 34% 5,7-dichloro-8-hydroxyquinoline-2-carboxylic acid [2-(1H-imidazol-4-yl)ethyl]amide (PBT 1038). This inhibited metal-mediated lipoprotein oxidation with IC50 value of 0.26 $\mu \rm M$.

MSTR 1

G14 = CONH2 (opt. substd.) / 62

Page 8

Patent location: claim 1

Note: also incorporates broader disclosure

Note: or pharmaceutically acceptable salts, hydrates, or

solvates

L5 ANSWER 2 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 149:121791 MARPAT

TITLE: Sox peptide-based sensor for detecting protein kinase

activity using chelation-enhanced fluorescence

INVENTOR(S): Imperiali, Barbara; Lukovic, Elvedin; Carrico-Moniz,

Dora

PATENT ASSIGNEE(S): Massachusetts Institute of Technology, USA

SOURCE: PCT Int. Appl., 36pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

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APPLICATION NO. DATE
PATENT NO. KIND DATE
_____
                                       _____
                                  WO 2007-US76959 20070828
    2008082715 A2 20080710 WO 2007-US76959 20070828
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA,
WO 2008082715
        CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI,
        GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG,
        KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL,
        PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN,
        TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
    RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
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        BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW,
        GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
        BY, KG, KZ, MD, RU, TJ, TM
                                        US 2006-511050
US 20080050761
                 A1 20080228
                                                         20060828
                                        US 2006-511050 20060828
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PRIORITY APPLN. INFO.:

US 2006-511050 20060828

AB The present invention provides sensors to monitor protein kinase activity continuously with a fluorescent read-out. The invention provides

continuously with a fluorescent read-out. The invention provides metal-binding compds. (Sox peptide) that exhibit chelation-enhanced fluorescence upon binding to Mg2+. The invention further provide peptidyl sensors which include a metal-binding peptide and one or two kinase recognition sequences with a hydroxyamino acid that can be phosphorylated in the presence of a kinase. The sensor peptides are synthesized via standard solid-phase peptide synthesis based on the optimal PKC peptide substrate and the activities of PKC isoenzymes were determined

MSTR 1

G5 = 72

G12-G13

—G6 G12—G13

= 120 / 125 G8

G12-G13 C(0)-G16

G12 = NH

G13 = cycloalkyl <containing 3-6 C> (opt. substd.)

G16 = NH2 / 92

9215—G13

Patent location: claim 1

Note: substitution is restricted

L5 ANSWER 3 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 149:104680 MARPAT

TITLE: Novel thiazolidine compounds as cannabinoid receptor

ligands and uses thereof

Carroll, William A.; Dart, Michael J.; Li, Tongmei; INVENTOR(S):

Perez-Medrano, Arturo V.; Peddi, Sridhar

PATENT ASSIGNEE(S): Abbott Laboratories, USA U.S. Pat. Appl. Publ., 40pp. SOURCE:

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE GΙ

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US 2007-954956
WO 2007-119974
     US 20080153883 A1 20080626
WO 2008079687 A1 20080703
                                                             20071212
                                            WO 2007-US87175 20071212
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA,
             CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI,
             GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG,
             KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME,
             MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL,
             PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN,
             TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW,
             GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
             BY, KG, KZ, MD, RU, TJ, TM
PRIORITY APPLN. INFO.:
                                            US 2006-876604P 20061222
```

$$\begin{array}{c|c}
 & R6 & O \\
 & R5 & X & || & \\
 & R4 & N & R1 \\
 & R3 & R2 & R2
\end{array}$$

The present invention relates to thiazolidinylidene containing compds. I [R1 = AΒ Ph (substituted with 1 to 5 Rj), naphthyl, cycloalkly, heterocyclyl, 2-Rg-pyridin-3-yl, quinolin-8-yl, benzofuran-5-yl, benzothien-5-yl; R2 =alkyl, alkoxy-(C2-6-alkylene), alkoxyalkoxy-(C2-6-alkylene), alkenyl, alkynyl, arylalkyl, cycloalklyalkyl, cycloalkoxyalkyl, (cycloalkylalkoxy)alkyl, cyanoalkyl, nitroalkyl, haloalkyl, haloalkoxyalkyl, heteroarylalkyl, heterocycloalkyl, (heterocyclyloxy)alkyl, hydroxyalkyl, etc.; R3, R4 = H, alkyl, cycloalkyl, haloalkly, heterocyclyl, hydroxyalkyl; CR3R4 = monocyclic cycloalkyl or heterocyclic ring, whereby the heterocycle contains at least one oxygen; R5, R6 = H, alkyl, aryl, cycloalkyl, haloalkly, heteroaryl, heterocyclyl, hydroxyalkyl; CR5R6 = monocyclic cycloalkyl or heterocyclic ring; R3C-CR5 = monocyclic cycloalkyl or heterocyclyl ring provided that the heterocycle is saturated and contains at least one oxygen; Rj, Rg = alkenyl, alkoxy, alkoxyalkoxy, alkoxyalkoxyalkyl, alkoxyalkyl, alkoxycarbonyl, alkoxycarbonylalkyl, etc.; X = S, 0] or a pharmaceutically acceptable salt thereof, compns. comprising such compds., and methods of treating conditions and disorders using such compds. and compns. Thus, thiazolidinylidene (Z)-I [R1 = 2-methoxy-5-chlorophenyl; R2 = CH2CH2OMe, R3 = R4 = H, R5 = R6 = Me; X = S] was prepared from 5-C1-2-MeOC6H4CO2H via amidation with 5,5-dimethyl-4,5-dihydro-1,3-thiazolyl-2-amine hydrochloride in THF containing HOBt and Et3N and N-alkylation with BrCH2CH2OMe in THF/DMF containing NaH. The cannabinoid receptor activity of thiazolidinylidenes I was tested [Ki < 1000 nM vs. CB2 receptor and Ki =10 to 1000 fold higher vs. CB1 receptor].

MSTR 1

G1 = 23

G4 = alkylcarbonyl < containing 1-10 C>

G5 = NH2

G24 = 109 / 114

HN—G4 C(O)-G5

Patent location: claim 1

Note: or pharmaceutically acceptable salts

L5 ANSWER 4 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 149:17765 MARPAT

ACCESSION NORDER. 149.17705 MAREAI

TITLE: Controlled-release formulation of piperazine-piperidine antagonists and agonists of the 5-HT1A

receptor having enhanced intestinal dissolution
INVENTOR(S): Ku, Mannching Sherry; Dulin, Wendy Ann; Lin, Yanning

Angela

PATENT ASSIGNEE(S): Wyeth, John, and Brother Ltd., USA

SOURCE: PCT Int. Appl., 91pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.		KI	ND :	DATE			A.	PPLI	CATI	ON N	٥.	DATE			
WO 2008	0673	 99	A	2 .	2008	0605		M	0 20	07-U	 S857	90	2007	1128		
W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,
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	GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,
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	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW				
RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,

IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

US 20080199518 A1 20080821 US 2007-986991 20071127 PRIORITY APPLN. INFO.: US 2006-861409P 20061128

AB The present invention relates to controlled-release beads comprising diquinoline-substituted piperazine-piperidine compds., such as 5-fluoro-8-[4-[4-(6-methoxyquinolin-8-yl)piperazin-l-yl]piperidin-l-yl]quinoline, or pharmaceutically acceptable salts thereof; to multiple particulate formulations comprising such beads; to methods of preparing such beads; and to methods of treating 5-HT1A-related disorders using such beads and/or multiple particulate formulations. Thus, beads were prepared containing sugar spheres coated with 5-fluoro-8-[4-[4-(6-methoxyquinolin-8-yl)piperazin-1-yl]piperidin-1-yl]quinoline trisuccinate, Opadry Clear II, and Surelease with or without citric acid. The dissoln. of active agent was enhanced in the presence of citric acid.

MSTR 1

G1 = 51 / 64

G3—G4 G8—G5 51 64

G3 = NH

G4 = alkyl < containing 1-6 C >

(opt. substd. by 1 or more G2)

G5 = NH2 / 66

66—G4

G8 = SO2 / C(O)

Patent location: claim 1

Note: or pharmaceutically acceptable salts

L5 ANSWER 5 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 148:538091 MARPAT

TITLE: Preparation of quinoline carboxamides as CSF 1R kinase

inhibitors for treating cancer and other diseases

INVENTOR(S): Dakin, Leslie; Daly, Kevin; Del Valle, David; Gero,

Thomas; Ogoe, Claude Afona; Scott, David; Zheng,

Xiaolan

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca Uk Limited

SOURCE: PCT Int. Appl., 84pp., which

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT	NO.		KI	ND	DATE			A.	PPLI	CATI	и ис	Э.	DATE			
WC	2008	0561	48	 A	 1	2008	0515		M	0 20	 07-G:	 В426.	3	2007	1108		
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,
		CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,	ES,	FI,
		GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,
		KM,	KN,	KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	ME,
		MG,	MK,	MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NΖ,	OM,	PG,	PH,	PL,
		PT,	RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ΤJ,	TM,	TN,
		TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW				
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,
		IS,	IT,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG,	BW,
		GH,	GM,	ΚE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,
		BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM									
PRIORIT	Y APP	LN.	INFO	.:					U	S 20	06-8	6524	5P	2006	1110		
									U	S 20	07-9	1618.	2P	2007	0504		

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention relates to chemical compds. of formula I (wherein one of R1 and R2 is selected from C1-6alkyl, C2-6alkenyl, etc. and the other R1 or R2 is H, halo, etc.; R3 is H or halo; R4 is halo, nitro, cyano, etc.; and n = 0-3) or pharmaceutically acceptable salts thereof which possess CSF 1R kinase inhibitory activity and are accordingly useful for their anticancer activity and thus in methods of treatment of the human or animal body. The invention also relates to processes for the manufacture of said chemical compds., to pharmaceutical compns. containing them and to their use in the manufacture of medicaments of use in the production of an anti cancer effect

in a warm blooded animal such as man. Example compound II, prepared from the corresponding tert-Bu carbamate III (preparation given), had an IC50 of 0.002 μM in an in vitro AlphaScreen assay that measures phosphorylation of a CSF-1R substrate.

MSTR 1A

GΙ

G1 = carbon chain <containing 1-6 C,

0 or more double bonds, 0 or more triple bonds>

(opt. substd. by G5)

G2 = 113

$$G10 = 313$$

$$G26 = 77-69 78-80 81-89$$

G30 = NH2

G34 = NH2

G35 = 90

 $G36 = 318-108 \ 320-314 \ 321-315 \ 316-226$



Patent location: claim 1

Note: substitution is restricted

Note: S-oxides

Note: or pharmaceutically acceptable salts

Note: also incorporates claim 8, formulas IV, V, VI,

VIIa, and VIIb

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 6 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 148:278889 MARPAT

TITLE: Sox peptide-based sensor for detecting protein kinase

activity using chelation-enhanced fluorescence

INVENTOR(S): Imperiali, Barbara; Lukovic, Elvedin; Carrico-Moniz,

Dora

PATENT ASSIGNEE(S): Massachusetts Institute of Technology, USA

SOURCE: U.S. Pat. Appl. Publ., 17pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO. DATE
US 20080050761 WO 2008082715	A1 2008022 A2 2008071	
W: AE, AG, CH, CN, GB, GD, KM, KN, MG, MK, PT, RO,	AL, AM, AT, AU CO, CR, CU, CZ GE, GH, GM, GT KP, KR, KZ, LA MN, MW, MX, MY RS, RU, SC, SD	, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN,
RW: AT, BE, IS, IT, BJ, CF, GH, GM,	BG, CH, CY, CZ LT, LU, LV, MC CG, CI, CM, GA	, UZ, VC, VN, ZA, ZM, ZW , DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, , MT, NL, PL, PT, RO, SE, SI, SK, TR, BF, , GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, , NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, , TM

PRIORITY APPLN. INFO.: US 2006-511050 20060828

AB The present invention provides sensors to monitor protein kinase activity continuously with a fluorescent read-out. The invention provides metal-binding compds. (Sox peptide) that exhibit chelation-enhanced fluorescence upon binding to Mg2+. The invention further provide peptidyl sensors which include a metal-binding peptide and at least one kinase recognition sequence with a hydroxyamino acid that can be phosphorylated in the presence of a kinase. The sensor peptides are synthesized via standard solid-phase peptide synthesis based on the optimal PKC peptide substrate and the activities of PKC isoenzymes were determined

MSTR 1

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10/572,914
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G5 = 72

G12—G13

G6 = NH2

G7 = 30 / 144

02S----G6 30 G12-G13

G8 = 120 / 125

G12-G13 125(0)-G16

G12 = NH

= cycloalky: = NH2 / 92 G13 = cycloalkyl <containing 3-6 C> (opt. substd.)

G16

G15—G13

Patent location: claim 1

Note: substitution is restricted

ANSWER 7 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 148:232646 MARPAT

TITLE: Fluorogenic protein kinase peptide substrates

comprising a fluorophore conjugated to a chelator

INVENTOR(S): Gee, Kyle

PATENT ASSIGNEE(S): Invitrogen Corporation, USA

SOURCE: PCT Int. Appl., 52pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.		KII	ND	DATE			A)	PPLI	CATI	ON No	Э.	DATE			
							_								
WO 20080167	62	A.	1	2008	0207		Mo	20	07-U	S730	0 0	2007	0706		
W: AE,	AG,	AL,	ΑM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,
CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,	ES,	FI,
GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,
KM,	KN,	KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	ME,
MG,	MK,	MN,	MW,	MX,	MY,	MZ,	NA,	NG,	ΝI,	NO,	NΖ,	OM,	PG,	PH,	PL,
PT,	RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ТJ,	TM,	TN,
TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW				
RW: AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
IS,	ΙΤ,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,

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BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

US 20070196860 A1 20070823 US 2007-624686 20070118

US 20080009026 A1 20080110 US 2007-774554 20070706

PRIORITY APPLN. INFO.: US 2006-819432P 20060707
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US 2007-624686

US 2006-759919P 20060118

20070118

AB The present invention relates to protein kinase sensors comprising a metal-chelating quinoline attached to a fluorophore and an amino acid. The invention also relates to methods of using these protein kinase sensors as well as kits comprising the protein kinase sensors.

MSTR 1

G2 = 76

G3-G11

G3 = NH (opt. substd.) G5 = 78 / CONH2 (opt. substd.) / SO2NH2 (opt. substd.)

G3-G11

G11 = acyl

Patent location: claim 8

Note: or tautomers, or salts

Stereochemistry: or stereoisomers

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 8 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 148:191837 MARPAT

TITLE: 3-Azabicyclo[3.1.0]hexane derivatives as vanilloid

receptor ligands, pharmaceutical compositions

containing them and process for their preparation

INVENTOR(S): Gharat, Laxmikant Atmaram; Joshi, Neelima Khairatkar;

Gajera, Jitendra Maganbhai; Yadav, Pravin Sabhajit

PATENT ASSIGNEE(S): Glenmark Pharmaceuticals S.A., Switz.

SOURCE: PCT Int. Appl., 116pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATEN	I NO.		KI	ND				A	PPLI	CATI	ON N	Ο.	DATE			
	 080100 080100							M	0 20	07-I	B200	2	2007	0716		
	: AE, CH, GB, KM, MG, PT,	AM, CR, GH, KR, MW, RU,	AT, CU, GM, KZ, MX, SC,	AU, CZ, GT, LA, MY, SD,	DE, HN, LC, MZ, SE,	DK, HR, LK, NA, SG,	DM, HU, LR, NG, SK,	DO, ID, LS, NI, SL,	DZ, IL, LT, NO, SM,	EC, IN, LU, NZ, SV,	BW, EE, IS, LY, OM, SY,	EG, JP, MA, PG,	ES, KE, MD, PH,	FI, KG, ME, PL,		
R	W: AT, IS, BJ, GH,	IT, CF,	BG, LT, CG, KE,	CH, LU, CI, LS,	CY, LV, CM,	CZ, MC, GA, MZ,	DE, MT, GN, NA,	DK, NL, GQ, SD,	EE, PL, GW, SL,	ES, PT, ML, SZ,	FI, RO, MR, TZ,	FR, SE, NE,	GB, SI, SN, ZM,	SK, TD,	TR, TG,	BF, BW,
PRIORITY A	,	,	,	,	1.0,	-0,	,	U U U	N 20 S 20 N 20 S 20	06-M 06-8 07-M 07-8	U113 3556 U381 9367	0P 5P	2006 2006 2007 2007 2007	0803 0227 0308		

GΙ

AB The invention relates to substituted 3-azabicyclo[3.1.0]hexane derivs., which are useful as vanilloid receptor ligands, methods of treating

diseases, conditions and/or disorders modulated by vanilloid receptors with them, and processes for preparing them. Compds. of formula I wherein X is O and S; R1 is quinolinyl, isoquinolinyl, 2-oxodihydroquinolinyl, and 1-oxodihydroisoquinolinyl; R2 and R3 are independently H, OH, and C1-6 alkyl; R4 and R5 are independently H, halo and alkyl; R4R5 taken together to form =O and =S; R6 is H, NO2, CN, CHO, Ac, halo, OH and derivs., SH and derivs., (un)substituted alkyl, (un)substituted (hetero)aryl, etc.; and their prodrugs, pharmaceutically acceptable salts, N-oxides, esters, solvates, tautomers, stereoisomers and polymorphs thereof, are claimed. Example compound II was prepared by a general procedure (procedure given). All the invention compds. were evaluated for their TRPV1 inhibitory activity (data given).

MSTR 1

$$\begin{array}{c|c} & G19 \\ G19 & 52 \\ 53 & G18 - G28 \\ G19 & 10 \\ 54 & G19 \\ & 55 \end{array}$$

$$G4 = NH$$
 $G6 = 20 / 46$

$$G8 = 39$$

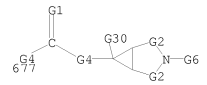
$$G14 = C(0) / S02$$

 $G18 = 102-10 \ 101-52 \ 96-53 \ 97-54 \ 98-55$

G19 = 403

G20 = NH2 (opt. substd.) / heterocycle <containing 3-7
 atoms, 1 or more heteroatoms, 1 or more N, zero or more O,
 zero or more S (no other heteroatoms),
 attached through 1 or more N> (opt. substd.)

G28 = 677



Patent location: claim 1

Note: additional derivatization also claimed

Note: or prodrugs, pharmaceutically acceptable salts, N-oxides, esters, solvates, tautomers or polymorphs Note:

also incorporates claim 43, structure 7 and claim

46, structure 8b

or stereoisomers Stereochemistry:

ANSWER 9 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

148:93193 MARPAT ACCESSION NUMBER:

TITLE: Method using fused heterocyclic compounds for the

treatment of glioma brain tumors

Bush, Ashley INVENTOR(S):

Prana Biotechnology Limited, Australia PATENT ASSIGNEE(S):

SOURCE: PCT Int. Appl., 115pp.

CODEN: PIXXD2

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE MO 2007145235 _____ WO 2007147217 A1 20071227 WO 2007-AU876 20070622 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO.: US 2006-815779P 20060622

The invention discloses therapeutic agents, formulations comprising them, and their use in the treatment, amelioration and/or prophylaxis of glioma brain tumors and related conditions. The therapeutic agent comprises two fused 6-membered rings with at least a nitrogen at position 1 and a hydroxyl at position 8.

MSTR 1

G1 = 33

C----G5

 $G2 = 20-2 \ 19-4 \ 23-37$

G4 = 38 / 50

G7—G8 G12—G13

G5 = SO2NH2 (opt. substd.)

G6 = 35

35—G4

G7 = NH

G8 = cycloalkyl <containing 3-6 C> (opt. substd.)

G12 = C(0)

G13 = NH2 / 52

G14-G8

Patent location: claim 1

Note: substitution is restricted

Note: and salts, hydrates, solvates, derivatives,

prodrugs, tautomers and isomers

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 10 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 148:55103 MARPAT

TITLE: Process for preparation of 8-piperazinyl-quinoline

derivatives

INVENTOR(S): Liu, Weiguo; Dragan, Vladimir; Strong, Henry Lee; Wu,

Yanzhong; Wen, Zhixin; Liang, Jessica Kangping; Durutlic, Haris; Sutherland, Karen Wiggins; Pilcher,

Anthony Scott

PATENT ASSIGNEE(S): Wyeth, John, and Brother Ltd., USA

SOURCE: PCT Int. Appl., 98pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	PATENT NO.					DATE			Α.	PPLI	CATI	N NC	Ο.	DATE			
WO	2007	1460	 72	 A	2	2007	1221		M	20 D	07-U	S134	 33	2007	0607		
WO	2007	1460	72	A	3	2008	0529										
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	ΒZ,	CA,
		CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,	ES,	FΙ,
		GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,
		KM,	KN,	KP,	KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	ME,
		MG,	MK,	MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,
		PT,	RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ΤJ,	TM,	TN,
		TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW				
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
		IS,	IT,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	ΤĠ,	BW,
														ZM,			
		BY,	KG,	KZ,	MD,	RU,	ΤJ,	TM,	AP,	EA,	EP,	OA	·	·			
US	2008	0058	523	Ā	1	2008	0306	·	Ü	s 20	07-8	1132	8	2007	0607		
PRIORIT	Y APP	LN.	INFO	. :					U	S 20	06-8	1214	8P	2006	0609		
OTHER SO	OURCE	(S):			CAS	REAC	T 14	8:55	103								

AB The present invention relates to processes for the preparation of 8-piperazinyl-quinoline derivs. with general formula I [wherein R1 - R6 = independently H, alkyl, alkenyl, halo, etc; R7 and R8 = independently H or CH3] or pharmaceutically acceptable salts thereof as 5-hydroxytryptamine receptor 1A (5-HT1A) binding agents, particularly as 5-HT1A receptor

antagonists or agonists. For example, 6-methoxy-8-(1-piperazinyl)quinoline (preparation given) was condensed with 1-(5-fluoroquinolin-8-yl)piperidin-4-one (preparation given) in presence of sodium triacetoxyborohydride in toluene at about 30 °C to give II as a product, which was further transformed to the tri-succinate salt thereof. Advantageously, the title processes allow for safer and environmentally tolerant production of these useful compds.

MSTR 1

G1 = 99 / 106 / 114

G5 = alkyl <containing 1-6 C>

(opt. substd. by 1 or more G4)

G6 = NH

G7 = NH2 / 108

168—G5

G9 = bond

G10 = NH2 / 132

132-G5

Patent location: claim 26

L5 ANSWER 11 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 148:54882 MARPAT

TITLE: Preparation of heteroaryl amides that interact with ion channels, in particular with ion channels from the

Kv family

INVENTOR(S): Blom, Petra; Defert, Olivier; Kaletta, Titus; Leysen,

Dirk Casimir Maria

PATENT ASSIGNEE(S): Devgen N.V., Belg. SOURCE: PCT Int. Appl., 62pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT NO. KIR				ND	DATE			A.	PPLI	CATI	и ис	Ο.	DATE			
WC	2007	1381	 12	 A	2	2007	 1206		M.	0 20	 07-е:	P554	08	2007	0601		
WC	2007	1381	12	A.	3	2008	0515										
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,
		CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,	ES,	FI,
		GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,
		KM,	KN,	KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	MG,
		MK,	MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NΙ,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,
		RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ТJ,	TM,	TN,	TR,
		TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	VN,	ZA,	ZM,	ZW					
	RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,
		IS,	ΙΤ,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG,	BW,
		GH,	GM,	ΚE,	LS,	MW,	MΖ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,
		BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM,	AP,	EA,	EP,	ΟA					
PRIORIT	Y APP	LN.	INFO	.:					E:	P 20	06 - 4	4707	5	2006	0601		
									U	S 20	06-8	0984	1P	2006	0601		

GΙ

$$\begin{bmatrix} \begin{bmatrix} \mathbf{R} \mathbf{1} \end{bmatrix}_{\mathbf{n}} & \begin{bmatrix} \mathbf{X}^2 \\ \mathbf{X}^1 & \mathbf{N} - \mathbf{L}^1 \end{bmatrix}_{\mathbf{R}} \end{bmatrix}_{\mathbf{R}} \begin{bmatrix} \mathbf{R}^2 \end{bmatrix}_{\mathbf{m}}$$

$$\begin{bmatrix} \mathbf{R} \mathbf{1} \end{bmatrix}_{\mathbf{n}} \begin{bmatrix} \mathbf{X}^3 \\ \mathbf{X}^4 \end{bmatrix} \begin{bmatrix} \mathbf{Z}^1 \\ \mathbf{N} \\ \mathbf{R}^3 \end{bmatrix} \begin{bmatrix} \mathbf{L}^1 \\ \mathbf{R}^2 \end{bmatrix}_{\mathbf{m}} \mathbf{I} \mathbf{I}$$

AB The present invention relates to compds. that interact with ion channels. In particular, the invention relates to compds. I or II [n, m = 0-4; Z1 = C(0), C(S), SO2; L1 = (un) substituted alkylene, cycloalkylene,

cycloalkylenoxyalkylene; X1 = O or S; X2 = CR4 or N; X3 = CR1 or N; X4 = CR1 or N; R1 = H, halo, OH, etc.; R2 = H, halo, OH, etc.; R3 = H, alkyl, aryl, etc.; R4 = H, halo, NH2, etc.; with the provisos]. Sixty-two specific title compds. such as III were prepared and/or claimed. The exemplified title compds. were tested in patch clamp assays (for example, III showed above 50% inhibition on Kv4.3-mediated potassium channel). The invention also relates to methods for preparing said compds. I (general protocols and schemes were given), to pharmaceutical compns. comprising said compds., and to the use of said compds. in methods for treatment of the human and animal body.

MSTR 1

$$G1 = 70$$

$$G8 = 38 / 40$$

G12 = NH2 G13 = NH

G14 = carbocycle <containing 3 or more C, non-aromatic,

0 or more double bonds, mono- or polycyclic> (opt. substd.)

G24 = 75 / N

Patent location: claim 1

Note: or tautomers, pharmaceutically acceptable salts or

solvates

Note: substitution is restricted Stereochemistry: or stereoisomers or racemics

L5 ANSWER 12 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 147:534639 MARPAT

TITLE: 3,4-Disubstituted coumarin and quinolone compounds for

the treatment of hepatitis C virus infection

INVENTOR(S): Xu, Bin; Zhu, Qiang; Cho, Hyun-Joon; Fathi, Reza;

Yang, Zhen; Sandrasagra, Anthony; Liu, Yixin

PATENT ASSIGNEE(S): XTL Biopharmaceuticals, Ltd., Israel

SOURCE: PCT Int. Appl., 150pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATEN'	I NO.		KI	ND	DATE			A	PPLI	CATI	ON N	0.	DATE			
WO 20	07133.	 211	 A	1	2007	 1122		W	0 20	 06-U	 S188	 57	2006	 0515		
W	: AE	, AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
	CN	, co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
	GE	, GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ΚM,	KN,	KP,	KR,
	ΚZ	, LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
	MZ	, NA,	NG,	ΝI,	NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
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	VN	, YU,	ZA,	ZM,	ZW											
R1	W: AT	, BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,
	IS	, IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
	CF	, CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG,	BW,	GH,
	GM	, KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
	KG	, KZ,	MD,	RU,	ТJ,	TM										

PRIORITY APPLN. INFO.:

WO 2006-US18857 20060515

AB The invention discloses 3,4-disubstituted coumarin and quinolone derivs. and processes for their preparation The invention also discloses methods for treating Hepatitis C virus infection by administering a 3,4-disubstituted coumarin or quinolone derivative

MSTR 1

G1 = 15

G2—G3

G2 = SO2

G3 = heteroaryl <containing zero or more N,

zero or more O, zero or more S> (opt. substd.)

G12 = 121

G13 = 83

c(0)-G16

G16 = NH2 G21 = NH G23 = 103

Patent location: claim 1

Note: or pharmaceutically acceptable salts or hydrates

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 13 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 147:480413 MARPAT

TITLE: Method using PB-1033 and related compounds for the

treatment of age-related macular degeneration (AMD)

INVENTOR(S): Bush, Ashley; Masters, Colin Louis PATENT ASSIGNEE(S): Prana Biotechnology Ltd, Australia

SOURCE: PCT Int. Appl., 109pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	ΓΕΝΤ	NO.		KI	ND	DATE			A	PPLI	CATI	ON N	Э.	DATE			
WO	2007	 1182	 76	 A	 1	2007	 1025		W	20 D	 07-A	 U490		 2007	0413		
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,
		CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,
		GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,
		KN,	KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	MG,	MK,
		MN,	MW,	MX,	MY,	MZ,	NA,	NG,	ΝI,	NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,
		RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ТJ,	TM,	TN,	TR,	TT,
		TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW						
	RW:	ΑT,	ΒE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,
		IS,	ΙT,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG,	BW,
		GH,	GM,	ΚE,	LS,	MW,	MΖ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,

BY, KG, KZ, MD, RU, TJ, TM

Ι

PRIORITY APPLN. INFO.:

US 2006-792278P 20060414

GΙ

AB The invention relates generally to the field of treatment and prophylaxis of retinal degenerative diseases. More particularly, the invention contemplates a method for preventing, reducing the risk of development of, or otherwise treating or ameliorating the symptoms of, age-related macular degeneration (AMD) or related retinal conditions in mammals and in particular humans. The invention further provides therapeutic compns. enabling dose-dependent or dose-specific administration of agents useful in the treatment and prophylaxis of age-related macular degeneration or related retinal degenerative conditions. Compds. useful invention include PB-1033 (I) and related compds.

MSTR 1

G14 = CONH2 (opt. substd.) / 62

Patent location: disclosure

Note: or salts, hydrates, solvates, derivatives,

prodrugs, tautomers

Note: substitution is restricted

Stereochemistry: or isomers

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 14 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

147:469247 MARPAT ACCESSION NUMBER:

Preparation of quinolones derivatives useful as TITLE:

inducible nitric oxide synthase inhibitors

INVENTOR(S): Roppe, Jeffrey R.; Bonnefous, Celine; Smith, Nicholas

D.; Lindstrom, Andrew K.; Noble, Stewart A.; Hassig, Christian A.; Payne, Joseph E.; Zhuang, Hui; Chen,

Xiaohong; Duron, Sergio G.

PATENT ASSIGNEE(S): Kalypsys, Inc., USA SOURCE:

PCT Int. Appl., 238pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.					ND	DATE			A.	PPLI	CATI	и ис	Э.	DATE					
		2007		. •	A2					M	0 20	0223								
	WO	2007117778			A.	3	20080207													
		W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	ΒA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,		
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,		
			GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,		
			KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,		
			MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,		
			RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	TJ,	TM,	TN,	TR,	TT,		
			TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW								
		RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,		
			IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,		
			CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,		
			GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,		
			KG,	KΖ,	MD,	RU,	ΤJ,	TM,	AP,	EA,	EP,	OA	,	·	·	·	,	•		
										US 2007-678572 20070223										
PRIOF	RIORITY APPLN. INFO.:										-		7656		20060224					
											US 2006-848696P 20061002									

GΙ

$$\begin{bmatrix} A & R^1 \\ B & & & \\ C & & & \\ D & R^3 & I \end{bmatrix}$$

AΒ The invention relates to novel quinolones of formula I [R1 =(un) substituted acyl, alkyl, alkylene, aminoalkyl, amidoalkyl, alkynyl, aryl, arylalkyl, arylalkoxy, etc.; R2 = (un)substituted acyl, alkoxy,

alkoxyalkyl, alkyl, alkylene, alkylamino, alkynyl, alkylimino, etc.; R2 may combine with R1 to form (un) substituted heterocycloalkyl; R3 = H, NH2, (un) substituted aryl, haloalkyl, (hetero) arylalkyl, (hetero) (cyclo) alkyl; A, B, C and D independently = (un) substituted acyl, alkoxy, alkyl, alkylene, alkylamino, alkynyl, etc.; any two or more A, B, C and D may combine to form aryl, cycloalkyl, heteroaryl or heterocycloalkyl], and their pharmaceutically acceptable salts, esters or prodrugs, are prepared and disclosed as inducible nitric oxide synthase (iNOS) inhibitors. Thus, e.g. II was prepared by acylation of aniline with Et 3-oxobutanoate followed by bromination and cyclization to generate intermediate 4-(bromomethyl)quinolin-2(1H)-one, which underwent substitution with aniline and acylation with furan-2-carbonyl chloride to provide II. The inhibitory activity of all exemplary compds. was evaluated in DAN assay and II was found to have EC50 value of \leq 5 μM . I should prove useful for inhibiting or modulating nitric oxide synthase and/or lowering nitric oxide levels of iNOS and for the treatment of an iNOS-mediated disease in a patient in need thereof.

MSTR 1

G11 G1 G11 G7 G11 G9

G1 = heteroarylamino <containing 1 or more heteroatoms,

zero or more N, zero or more O,

zero or more S (no other heteroatoms)> (opt. substd.)

G7 = CONH2 (opt. substd.) G11 = SO2NH2 (opt. substd.)

Patent location: claim 1

Note: or salts, esters or prodrugs

Note: additional substitution and ring formation also

claimed

L5 ANSWER 15 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 147:406803 MARPAT

TITLE: Preparation of benzenediamine derivatives as

inhibitors of the interactions between MDM2 and p53 INVENTOR(S): Lacrampe, Jean Fernand Armand; Meyer, Christophe;

Schoentjes, Bruno; Lardeau, Delphine Yvonne Raymonde;

Poncelet, Alain Philippe; Van Hijfte, Luc

PATENT ASSIGNEE(S): Janssen Pharmaceutica N.V., Belg.

SOURCE: PCT Int. Appl., 60pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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_____
     WO 2007107543 A1 20070927
                                          WO 2007-EP52579 20070319
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA,
             CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB,
             GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM,
             KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK,
             MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO,
             RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT,
             TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW,
             GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
             BY, KG, KZ, MD, RU, TJ, TM
PRIORITY APPLN. INFO.:
                                           EP 2006-111531
                                                            20060322
                                           US 2006-784780P 20060322
GΙ
```

A—
$$(CH_2)_n$$
— $(NH)_p$ — $(CH_2)_m$ — $(CH_2)_s$ — NH — $(CH_2)_t$ — Z

H
N
NH
NH

AB The title compds. I [wherein m = 0-2; n = 0-4; p, s, t independently = 0 or 1; R1, R2 independently = H, halo, alkyl, etc.; A = (un)substituted Ph, pyridinyl, pyrrolyl, thiophenyl or furanyl; Z = certain (un)substituted nitrogen heteroaryl] and N-oxides, salts, or stereoisomers thereof are prepared as inhibitors of the interactions between MDM2 and p53. For example, compound II was prepared in a multi-step synthesis. I showed inhibitory effect in both p53 ELISA assay and cell proliferation assay. The invented compds. are useful for the treatment of disorders mediated by p53-MDM2 interactions.

MSTR 1

G12 = NHG13 = 271

G16 = CONH2 G25 = 2-1 3-4

g2—g26

G26 = phenylene (opt. substd. by (1-2) G7)

Patent location: claim 1

Note: or N-oxides, addition salts

Note: also incorporates claim 10, formula (VIII)

Stereochemistry: or stereochemically isomeric forms

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 16 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 147:343961 MARPAT

TITLE: Quinoline derivatives as platelet-derived growth factor inhibitors, their preparation, pharmaceutical

compositions, and use in the treatment of cancer

compositions, and use in the treatment

INVENTOR(S):
Jung, Frederic Henri; Ple, Patrick

PATENT ASSIGNEE(S): AstraZeneca AB, Swed.; AstraZeneca UK Limited

SOURCE: PCT Int. Appl., 133pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	ENT 1	NO.		KIND DATE				APPLICATION NO. DATE										
WO 2007099335			 A	 1	2007	0907		WO 2007-GB728 20070301										
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KN,	
		KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	MG,	MK,	MN,	
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		IS,	ΙΤ,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	
		ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	${ m ML}$,	MR,	ΝE,	SN,	TD,	ΤG,	BW,	
		GH,	GM,	KΕ,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	
		BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM										

PRIORITY APPLN. INFO.:

EP 2006-300186 20060302 EP 2006-301104 20061031

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention concerns quinoline derivs. of formula I, which are inhibitors of platelet-derived growth factor (PDGF). In compds. I, p is 0-3; each R1 is independently halo, OH, NH2, SH, CF3, cyano, carboxy, C1-6 alkoxycarbonyl, carbamoyl, C1-8 alkyl, C1-6 alkoxy, C1-6 alkylthio, C1-6 alkylsulfonyl, C1-6 alkylamino, N-(C1-6 alkyl)carbamoyl, C2-6 acyl, C2-6 acylamino, etc.; q is 0, 1, or 2; each R2 is independently halo, OH, NH2, CF3, cyano, C1-8 alkyl, C2-8 alkenyl, C2-8 alkynyl, C1-6 alkoxy, C1-6 alkylamino, or di(C1-6 alkyl)amino; R3 is H, C1-8 alkyl, C2-8 alkenyl, or C2-8 alkynyl; R4 is H, C1-8 alkyl, C2-8 alkenyl, C2-8 alkynyl, C1-6 haloalkyl, C1-6 alkoxy-C1-6 alkyl, carboxy-C1-6 alkyl, etc., or R3 and R4, together with the carbon atom to which they are attached, form C3-8 cycloalkyl; R5 is H, C1-8 alkyl, C2-8 alkenyl, C2-8 alkynyl, or -L-R6; R6 is C1-6 haloalkyl, C1-6 hydroxyalkyl, C1-6 alkoxy-C1-6 alkyl, or cyano-C1-6 alkyl; L is a bond, O, or NR7, where R7 is H or C1-8 alkyl; X is O or NR8, where R8 is H or C1-8 alkyl; and A is (un)substituted aryl or (un) substituted heteroaryl containing up to three heteroatoms selected from O, N, and S. The invention also relates to the preparation of I, pharmaceutical compns. comprising a quinoline of formula I in association with a pharmaceutically acceptable diluent or carrier, as well as to the use of the compns. for the treatment of cell proliferative disorders, such as cancer. Deprotonation of acetonitrile and condensation with Et propionate gave 3-oxopentanenitrile, which underwent heterocyclocondensation with hydrazine to form 5-amino-3-ethylpyrazole (II). Hydrogenation of Et 2-(5-benzyloxypyrimidin-2-yl)acetate followed by substitution of 4-chloro-6,7-dimethoxyquinoline resulted in the formation of quinoline III, which was hydrolyzed and amidated with II to give quinoline IV. The compds. of the invention are inhibitors of PDGF, e.g., compound IV expressed IC50 value of 6 nM vs. phospho-Tyr751 formation in PDGFR β .

MSTR 1

G1 = CONH2 / alkylaminosulfonyl < containing 1-6 C> G6 = NH

Patent location: claim 1

Note: or pharmaceutically acceptable salts, solvates or

prodrugs

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 17 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 147:343960 MARPAT

TITLE: Quinoline derivatives as platelet-derived growth

factor inhibitors, their preparation, pharmaceutical

compositions, and use in the treatment of cancer

INVENTOR(S): Jung, Frederic Henri; Ple, Patrick

PATENT ASSIGNEE(S): AstraZeneca AB, Swed.; AstraZeneca UK Limited

SOURCE: PCT Int. Appl., 217pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

F	PATE	TENT NO.				KIND DATE						CATI	N NC	Э.	DATE				
 W	vo 2	2007	 0993:	 26	A1 20070907					M.	0 20	 07-G:	 В719		20070301				
		W: AE, AG,			AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
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			KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	MG,	MK,	MN,	
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			RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	
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			IS,	ΙΤ,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	
			ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	$\mathrm{ML}_{,}$	MR,	ΝE,	SN,	TD,	TG,	BW,	
			GH,	GM,	ΚE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,	
			BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM										
PRIORI	PRIORITY APPLN. INFO.:									EP 2006-300181 20060302									
										EP 2006-301102 20061031									

GI

The invention concerns quinoline derivs. of formula I, which are inhibitors of platelet-derived growth factor (PDGF). In compds. I, p is 0-3; each R1 is independently halo, OH, NH2, SH, CF3, cyano, carboxy, C1-6 alkoxycarbonyl, carbamoyl, C1-8 alkyl, C1-6 alkoxy, C1-6 alkylthio, C1-6 alkylsulfonyl, C1-6 alkylamino, N-(C1-6 alkyl)carbamoyl, C2-6 acyl, C2-6 acylamino, etc.; q is 0, 1, or 2; each R2 is independently halo, OH, NH2, CF3, cyano, carboxy, carbamoyl, C1-8 alkyl, C1-6 alkoxy, C1-6 alkylamino, N-(C1-6 alkyl)carbamoyl, C1-6 haloalkyl, C1-6 alkoxy-C1-6 alkyl, etc.; R3 is H, C1-8 alkyl, C2-8 alkenyl, or C2-8 alkynyl; R4 is H, C1-8 alkyl, C2-8 alkenyl, C2-8 alkyl, etc., or R3 and R4, together with the carbon atom to which they are attached, form C3-8 cycloalkyl; R5 is H, C1-8 alkyl, C2-8

^{*} STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

alkenyl, C2-8 alkynyl, or -L-R6; R6 is C1-6 haloalkyl, C1-6 hydroxyalkyl, C1-6 alkoxy-C1-6 alkyl, or cyano-C1-6 alkyl; L is a bond, O, or NR7, where R7 is H or C1-8 alkyl; X is O or NR8, where R8 is H or C1-8 alkyl; and A is (un)substituted aryl or (un)substituted heteroaryl containing up to three heteroatoms selected from O, N, and S. The invention also relates to the preparation of I, pharmaceutical compns. comprising a quinoline of formula I in association with a pharmaceutically acceptable diluent or carrier, optionally containing an addnl. antitumor or antiangiogenic agent, as well as to the use of the compns. for the treatment of cell proliferative disorders, such as cancer. N-Nitration of pyrazole followed by rearrangement gave 4-nitropyrazole, which was N-alkylated with di-Et sulfate and reduced to give 4-amino-1-ethylpyrazole (II). Substitution of 4-chloro-6-cyano-7methoxyquinoline with 2-(4-hydroxyphenyl)acetic acid yielded quinoline III, which underwent amidation with II to give quinoline IV. The compds. of the invention are inhibitors of PDGF, e.g., compound IV expressed IC50 value of 2 nM vs. phospho-Tyr751 formation in PDGFR β .

MSTR 1

G1 = CONH2 / alkylaminosulfonyl <containing 1-6 C>

G6 = NH

INVENTOR(S):

Patent location: claim 1

Note: or pharmaceutically acceptable salts, solvates or

prodrugs

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 18 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 147:322860 MARPAT

TITLE: Quinoline derivatives as platelet-derived growth

factor inhibitors, their preparation, pharmaceutical

compositions, and use in the treatment of cancer Jung, Frederic Henri; Morgentin, Remy Robert; Ple,

Patrick

PATENT ASSIGNEE(S): Astrazeneca A/B, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 155pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO. KIND DATE
                                         APPLICATION NO. DATE
    _____
                                        _____
    WO 2007099323 A2 20070907
WO 2007099323 A3 20071115
                                        WO 2007-GB713 20070301
           AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
            CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
            GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN,
            KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, MN,
            MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS,
            RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ,
            UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
        RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
            IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF,
            BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW,
            GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
            BY, KG, KZ, MD, RU, TJ, TM, AP, EA, EP, OA
                                        EP 2006-300183
PRIORITY APPLN. INFO.:
                                                          20060302
                                         EP 2006-301103 20061031
GΙ
```

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The invention concerns quinoline derivs. of formula I, which are inhibitors of platelet-derived growth factor (PDGF). In compds. I, p is 0-3; each R1 is independently halo, OH, NH2, SH, CF3, cyano, carboxy, C1-6 alkoxycarbonyl, carbamoyl, C1-8 alkyl, C1-6 alkoxy, C1-6 alkylthio, C1-6 alkylsulfonyl, C1-6 alkylamino, N-(C1-6 alkyl)carbamoyl, C2-6 acyl, C2-6 acylamino, etc.; q is 0, 1, or 2; each R2 is independently halo, OH, NH2, CF3, cyano, carboxy, carbamoyl, C1-8 alkyl, C1-6 alkoxy, C1-6 alkylamino, N-(C1-6 alkyl)carbamoyl, C1-6 haloalkyl, C1-6 alkoxy-C1-6 alkyl, etc.; R3 is H, C1-8 alkyl, C2-8 alkenyl, or C2-8 alkynyl; R4 is H, C1-8 alkyl, C2-8 alkenyl, C2-8 alkynyl, C1-6 haloalkyl, C1-6 alkoxyalkyl, carboxy-C1-6 alkyl, etc., or R3 and R4, together with the carbon atom to which they are attached, form C3-8 cycloalkyl; R5 is H, C1-8 alkyl, C2-8 alkenyl, C2-8 alkynyl, or -L-R6; R6 is C1-6 haloalkyl, C1-6 hydroxyalkyl, C1-6 alkoxy-C1-6 alkyl, or cyano-C1-6 alkyl; L is a bond, O, or NR7, where R7 is H or C1-8 alkyl; X is O or NR8, where R8 is H or C1-8 alkyl; and A is (un) substituted aryl or (un) substituted heteroaryl containing up to three heteroatoms selected from O, N, and S. The invention also relates to the preparation of I, pharmaceutical compns. comprising a quinoline of formula I in association with a pharmaceutically acceptable diluent or carrier, optionally containing an addnl. antitumor or antiangiogenic agent, as well as to the use of the compns. for the treatment of cell proliferative disorders, such as cancer. Benzylation of 5-hydroxy-2-methylpyridine followed by N-oxidation, acetylation, rearrangement, and hydrolysis gave pyridine II, which was chlorinated, substituted with cyanide, and hydrolyzed resulting in the formation of 2-pyridineacetic acid III. tert-Bu esterification of III, hydrogenation, and substitution of 4-chloro-6,7-dimethoxyquinoline yielded IV, which underwent acidic deesterification and amidation with 4-amino-1-ethylpyrazole (four-step preparation is given) to give quinoline V. The compds. of the invention, e.g., V, are PDGF inhibitors (no data).

MSTR 1

G1 = CONH2 / alkylaminosulfonyl <containing 1-6 C>

G6 = NH

Patent location: claim 1

Note: or pharmaceutically acceptable salts, solvates or

prodrugs

L5 ANSWER 19 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 147:160527 MARPAT

TITLE: Measuring protein kinase activity using

phosphorylatable peptides exhibiting increased

fluorescence when sensor moieties are complexed with

metal ions

INVENTOR(S): Schaefer, Erik M.; Qian, Xiao-Dong; Li, Min; Gee, Kyle

R.

PATENT ASSIGNEE(S): Invitrogen Corp., USA SOURCE: PCT Int. Appl., 78pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PAT	CENT :	NO.		KI	ND	DATE			Α.	PPLI	CATI	и ис	Э.	DATE			
WO	2007	0849	 68	A	1	2007	0726		M	20 C	 07-U	5607:	 29	2007	0118		
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
		GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KN,
		KP,	KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,
		MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,
		RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ΤJ,	TM,	TN,	TR,	TT,
		TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW						
	RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
		IS,	ΙT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG,	BW,	GH,
		GM,	KΕ,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,	BY,
		KG,	KΖ,	MD,	RU,	ТJ,	TM										
PRIORITY	APP	LN.	INFO	.:					U	S 20	06-7	5991	9P	2006	0118		
									U	S 20	06-8	1943	2P	2006	0707		

GΙ

The present invention relates to methods for detecting and/or measuring AΒ the activity of a specific protein kinase, with the methods comprising contacting one or more kinases with a binding agent to isolate a specific kinase of interest. The isolated kinase is then contacted with a kinase activity sensor, where the kinase activity sensor is comprised of a kinase recognition motif that is capable of being recognized by the isolated kinase, and at least one phosphorylation site. The isolated kinase phosphorylates the amino acid target of the kinase activity sensor and levels of the phosphorylated target amino acid can then be quantified. Thus, a mouse monoclonal antibody specific for p38 kinase is attached to the wells of a 96-well plate. After the antibody captures the specific kinase of interest (p38) from murine macrophage cells, a kinase activity sensor comprising the kinase recognition motif AHLQRLSI9(dP), where dP is D-proline, and the metal binding amino acid SOX (I) are added to the wells along with ATP. The SOX amino acid fluoresces upon chelation of the ternary complex with phosphorylated peptide and magnesium.

MSTR 2

$$G1 = 50 / 71$$

$$G4 = NH$$
 $G8 = NH2 / heterocycle < containing 1-4 heteroatoms,$

1 or more N, zero or more O, zero or more S (no other heteroatoms), 1-10 C, attached through 1 or more N>(opt. substd.)

G10 = 0

= 78 / SO2G14

_C===G10

G18 = 112 / 123 / 127

G4—C(0)-G2 G14—G8 G11—G15—G8

Patent location: claim 14

Note: or tautomers or salts or stereoisomers Stereochemistry:

5 REFERENCE COUNT: THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 20 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

146:337747 MARPAT ACCESSION NUMBER:

TITLE: Preparation of quinoline compounds as Met kinase

inhibitors for the treatment of cancer

INVENTOR(S): Kim, Kyoung S.

PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA

SOURCE: U.S. Pat. Appl. Publ., 22pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	CENT	NO.		KI	ND	DATE			A.	PPLI	CATI	ON N	Э.	DATE			
US	2007	0060	 613	 A	 1	2007			U	S 20	 06-5	2052	0	2006	 0913		
WO	2007	0331	96	A	1	2007	0322		M	O 20	06-U	S355.	28	2006	0913		
	W:	ΑE,	AG,	AL,	ΑM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
		GE,	GH,	GM,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	ΚM,	KN,	KP,
		KR,	KΖ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,
		MW,	MX,	MY,	MΖ,	NA,	NG,	ΝI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,
		RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	US,	UΖ,	VC,	VN,	ZA,	ZM,	ZW							
	RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,
		IS,	ΙΤ,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,
		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
		KG,	KΖ,	MD,	RU,	ТJ,	TM										
RITY	Z APP	LN.	TNFO	. •					IJ.	S 20	05 - 7	1686	4 P	2005	0914		

PRIORITY APPLN. INFO.:

US 2005-716864P 20050914

GΙ

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- AB Title compds. such as I [wherein B = 0, S, SO2, etc.; X, A, D = N or (un)substituted CH; R1 = H, halo, cyano, etc.; R3a, R4a, R9 = H, (un)substituted alkyl, aryl, etc.; R5 R8 = H, halo, NO2, etc.], which are useful as Met kinase inhibitors and anticancer agents (no data), were prepared For example, II was synthesized as TFA salt in 30% yield by amidation of the corresponding dihydropyridinecarboxylic acid with (quinolinyloxy)aniline.

MSTR 1

G1 = 194

G14-G13

G5 = 222 / 224 / 229 / 232

G14-G13 C(O)-G7 O2S-G13 G14-G9 222 224 229 232

G7 = 208

G14-G13

G13 = heteroaryl <containing 9-10 atoms, zero or more N,
 zero or more O, zero or more S (no other heteroatoms),
 bicyclic> (opt. substd.)

G14 = NHG17 = 183

C——G5

G18 = NH

Patent location: claim 1

L5 ANSWER 21 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 146:229194 MARPAT

TITLE: Preparation of polyguinoline metal ligand complexes

and the therapeutic use thereof in treatment of

neurodegenerative disorders

INVENTOR(S): Deraeve, Celine; Pitie, Marquerite; Boldron,

Christophe; Meunier, Bernard

PATENT ASSIGNEE(S): Palumed S.A., Fr.; Centre National De La Recherche

Scientifique (C.N.R.S)

SOURCE: PCT Int. Appl., 133pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PA:	CENT I	МО.		KI	ND	DATE			A.	PPLI	CATI	и ис	Э.	DATE			
		2007								W	0 20	06-F	R190	6	2006	0804		
	W O								Δ7.	RΔ	RR	BG	BB	ВW	BY,	B7.	$C\Delta$	СН
		VV •						•	•						ES,	•		
						•						•			KG,			
			•		•		•	•	•						MD,	•		•
			•	•	•	•	•	•		•	•	•	•	•	PT,	•	,	•
				•		•	•	•	•	•	•	•	•		TT,	•	,	•
				•	_ •	•	ZA,	•	•	,	,	,	,	•	•	,	•	,
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															SK,			
			CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,
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			KG,	KΖ,	MD,	RU,	ТJ,	TM,	AP,	EA,	EP,	OA						
	FR	2889	525		А	1	2007	0209		F	R 20	05-8	351		2005	0804		
	CA	2616	453		Α	1	2007	0208		C.	A 20	06-2	6164	53	2006	0804		
	EΡ	1919	894		A	2	2008	0514		\mathbf{E}	P 20	06 - 7	9429.	3	2006	0804		
		R:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,
			IS,	ΙT,	LI,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR	
PRIOR	IT	Y APP	LN.	INFO	.:					F.	R 20	05-8	351		2005	0804		
										W	0 20	06-F	R190	6	2006	0804		
GI																		

$$R^{2}$$
 R^{2}
 R^{3}
 R^{4}
 R^{5}
 R^{5}

Ι

II

AB Polyquinoline I, wherein X is OR, NRR', S(O)pR, OCOR, OCOOR, substituted

N-containing heterocycle; Y is N-containing heterocycle, H, OR, NRR', halogen, ${\rm CN}$,

CF3, alkyl; R and R' are independently H, cycloalkyl, alkyl; R1-R5 are independently H, OR, NRR', halogen, CN, CF3, S(0)pR, COOR, OCOOR, CONRR', NRCOOR', alkyl; p is 1-2; were prepared and used thereof in the form of therapeutic agents in treatment of neurodegenerative disorders such as Alzheimer, Parkinson, encephalopathy, Huntington, amyotrophic lateral sclerosis, Down's syndrome. Thus, quinoline II ligand complexes with copper and zinc were prepared and used in the treatment of neurodegenerative disorders. Title metal complexes were tested in vitro and used to dissolve β -amyloid peptide aggregates and inhibit or diminish to generation of H2O2 for the treatment of Alzheimer, Parkinson, encephalopathy, Huntington, amyotrophic lateral sclerosis, Down's syndrome diseases.

MSTR 1A

G7 = carbocycle <containing 3-11 C, non-aromatic, 0 or more double bonds, 1-3 rings> (opt. substd.)

G9 = NH2 / 46

$$G10 = 55 / 70$$

$$G11 = NH$$

 $G12 = 124$

$$G16 = 161$$

G11-G7

Patent location: claim 1

Note: substitution is restricted

Note: or pharmaceutically acceptable hydrates, solvates,

salts, or esters

Stereochemistry: or stereoisomers or mixtures

L5 ANSWER 22 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 146:100570 MARPAT

TITLE: Pyridinones and pyridazinones as potassium channel

inhibitors, their preparation, pharmaceutical

compositions, and use in therapy

INVENTOR(S): Brendel, Joachim; Englert, Heinrich Christian; Wirth,

Klaus; Wagner, Michael; Ruxer, Jean-Marie; Pilorge,

Fabienne

PATENT ASSIGNEE(S): Sanofi-Aventis Deutschland GmbH, Germany

SOURCE: PCT Int. Appl., 54pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO. KIND DATE
                                          APPLICATION NO. DATE
                                           _____
     WO 2006136304 A1 20061228 WO 2006-EP5578 20060610
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR,
             KZ, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX,
             MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD,
             SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ,
             VC, VN, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
             CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH,
             GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM
     DE 102005028862 A1 20070111
                                          DE 2005-10200502886220050622
                     A1
     AU 2006261316
                                           AU 2006-261316 20060610
                            20061228
     CA 2610075
                                           CA 2006-2610075 20060610
                      A1
                            20061228
                      A1
                                          EP 2006-754277 20060610
     EP 1896416
                           20080312
         R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR
                     A1 20080807
     US 20080188477
                                           US 2007-954396 20071212
     MX 200715970 A 20080306
IN 2007CN05882 A 20080627
KR 2008018903 A 20080628
CN 101203493 A 20080618
                                                             20071214
                                            MX 2007-15970
                                            IN 2007-CN5882
                                                              20071220
                                           KR 2007-730045 20071221
                                           CN 2006-80022542 20071221
PRIORITY APPLN. INFO.:
                                            DE 2005-10200502886220050622
                                            WO 2006-EP5578
                                                            20060610
```

GΙ

$$R^{1}$$
 R^{2}
 R^{5}
 R^{5}
 R^{5}
 R^{6}
 R^{6}

AΒ The invention relates to compds. of the general formula I, which are inhibitors of the Kv1.5 potassium channel. In compds. I, X is CH or N; R1 and R2 are independently selected from (un)substituted Ph, (un)substituted pyridinyl, (un)substituted thienyl, (un)substituted naphthyl, (un) substituted quinolinyl, (un) substituted pyrimidinyl, or (un)substituted pyrazinyl; R3 is (CH2)p-R7, where p is 0-5 and R7 is Me, CH2F, CHF2, CF3, C3-7 cycloalkyl, ethynyl, propynyl, C1-4 alkoxy, (un) substituted Ph, or (un) substituted 2-pyridinyl; R4 and R5 are independently selected from H and C1-3 alkyl; and R6 is H, F, C1, CF3, or C1-3 alkyl; including pharmaceutically acceptable salts thereof. The invention also relates to the preparation of I, pharmaceutical compns. comprising an effective amount of at least one compound I with pharmaceutically acceptable carriers and additives, optionally in combination with other pharmacol. active ingredients, as well as to the use of the compns. for the treatment and prophylaxis of atrial arrhythmias, for example atrial fibrillation (AF) or atrial flutter. Ring opening of racemic cis-stilbene oxide with 2(1H)-pyridinone followed by alkylation with cyclopropylmethyl bromide gave (R*,R*)-pyridinone II. Several compds. of the invention, e.g., II, express IC50 values for the Kv1.5 channel of less than 1 μ M.

MSTR 1

G2 = 234

= CONH2 / NMe2 / SO2NH2

G15 = N

Patent location: claim 1

Note: and pharmaceutically acceptable salts and

trifluoroacetates

Note: substitution is restricted

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 3 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 23 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER:

146:81895 MARPAT

TITLE:

Piperazine-piperidine antagonists and agonists of the 5-HT1A receptor and their preparation, pharmaceutical compositions, and use in the treatment of central

nervous system disorders

INVENTOR(S):

Asselin, Magda; Grosu, George Theodore; Sabb, Anmarie Louise; Childers, Wayne Everett; Havran, Lisa Marie; Shen, Zhongui; Bicksler, James Jacob; Chong, Dan

Chaekoo

PATENT ASSIGNEE(S): Wyeth, John, and Brother Ltd., USA

SOURCE:

PCT Int. Appl., 219pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	TENT	NO.		KII	ND	DATE			A.	PPLI	CATI	ON N	Ο.	DATE			
	2006 2006					 2006 2007			M.	0 20	 06-U	 S227	 19	2006	0609		
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SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC,
             VN, YU, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
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     AU 2006257874
                             20061221
                                            AU 2006-257874
                                                              20060609
                       Α1
     CA 2611711
                             20061221
                                            CA 2006-2611711
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                       Α1
     US 20070027160
                             20070201
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                       Α1
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     EP 1888559
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                                                              20060609
             AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL,
             BA, HR, MK, YU
     IN 2007KN04732
                                            IN 2007-KN4732
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                                                              20071210
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                                                              20071211
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PRIORITY APPLN. INFO.:
                                            US 2005-689469P
                                                              20050610
                                            WO 2006-US22719
                                                              20060609
GΙ
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AB The invention relates to novel piperazine-piperidine compds. of formula I. Compds. of formula I wherein each R are independently H, C1-6 (halo)alkyl, C2-6 alkenyl, C2-6 alkynyl, halo, CF3, NO2, CN, OH and derivs., OSO2H and derivs., SH and derivs., SO2H and derivs., etc.; each R' are independently H and Me; and their pharmaceutically acceptable salts are claimed. The compds. are useful as 5-HT1A binding agents, particularly as 5-HT1A receptor antagonists and agonists. These compds. are useful in treating central nervous system disorders, such as cognition disorders, anxiety disorders, depression and sexual dysfunction. Example compound II was prepared by cyclization of 4-amino-3-chlorophenol with glycerol; the

ΙI

resulting 8-chloro-6-hydroxyquinoline underwent methylation to give 8-chloro-6-methoxyquinoline, which underwent substitution with N-Boc-piperazine to give 6-methoxy-8-[1-(tert-butoxycarbonyl)-4-piperazino]quinoline, which underwent hydrolysis to give 6-methoxy-8-piperazinoquinoline, which underwent reductive alkylation with 1-(quinolin-8-yl)piperidin-4-one to give compound II. All the invention compds. were evaluated for their 5-HT1A antagonistic and agonistic activity. From the assay, it was determined that compound II exhibited an 5-HT1A

affinity with a Ki value of 0.40 nM and antagonistic activity with IC50 og 3.86 nM.

MSTR 1

G1 = 99 / 106 / 114

G6—G5 O2S—G7 G9—C(0)—G10

G2 = (0-2) CH2

G5 = carbon chain <containing 1-6 C,

0 or more double bonds, 0 or more triple bonds>

(opt. substd. by G4)

G6 = NH

G7 = NH2 / 108

G8---G5

G9 = bond

G10 = NH2 / 132

G12—G5

Patent location: claim 1

Note: substitution is restricted also incorporates claim 15

Note: and pharmaceutically acceptable salts and hydrates

L5 ANSWER 24 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 145:438538 MARPAT

TITLE: Preparation of quinolin-5-yl acylhydrazide derivatives

as p2x7 antagonists and use as antinociceptive

prodrugs

INVENTOR(S): Nelson, Derek W.; Jarvis, Michael F.; Carroll, William

Α.

PATENT ASSIGNEE(S): Abbott Laboratories, USA SOURCE: PCT Int. Appl., 79pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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US	2006	0276	505	A.	1	2006	1207		U	S 20	06 - 4	0049	2	2006	0407		
PRIORI	ORITY APPLN. INFO.:									S 20	05-6	7020	8P	2005	0411		
GI																	

AB Quinolin-5-yl acylhydrazide derivs. I wherein D is a 5 or 6 membered heteroaryl ring; A is an alkyl, cycloalkyl, heterocyclic ring, etc.; m is 0 to 3; n is 0 to 4; Rx and Ry are independently selected from alkyl, alkenyl, halo, nitro cyano, etc are prepared as prodrugs with antinociceptive properties. Thus, II was prepared and tested for its in vitro IL-1 β release and in vivo antinociceptive effects (no data). Further, I can be employed in the treatment of pain, neuropathic pain, inflammation, chronic inflammatory pain, neurodegeneration, depression and promoting neuroregeneration.

MSTR 1

10/572,914

$$G1 = 13$$

$$G2 = 15$$

$$G3 = 17$$

$$G4 = 22-2 23-5 19-14 20-16 21-18$$

$$G8 = NH$$

$$G8 = NH$$

 $G10 = alkylcarbonyl < containing 1-10 C>$
 $G14 = 150$

$$G14 = 150$$

$$G17 = 158$$

G8---G10

Patent location: claim 1

additional derivatization also disclosed Note: Note: additional oxo formation also claimed

Note: or pharmaceutically acceptable salts or prodrugs

REFERENCE COUNT: 11 THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 25 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 145:419178 MARPAT

TITLE: Preparation of novel substituted diazabicyclooctane

derivatives as monoamine neurotransmitter re-uptake

inhibitors

INVENTOR(S): Peters, Dan; Nielsen, Elsebet Oestergaard; Redrobe,

John Paul

PATENT ASSIGNEE(S): Neurosearch A/S, Den. PCT Int. Appl., 25pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

]	PATEN:	r no.		KI	ND	DATE			A.	PPLI	CATI	и ис	ο.	DATE			
Ţ	WO 200	061060	90	A	1	2006	1012		M	0 20	06-E	P612	61	2006	0403		
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		KG,	KΖ,	MD,	RU,	ТJ,	TM										
]	EP 186	59050		A	1	2007	1226		E:	P 20	06-7	2550	7	2006	0403		
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PRIOR	ITY A	PPLN.	INFO	.:					D:	K 20	05 - 4	66		2005	0404		
									U	S 20	05-6	6766	9P	2005	0404		
									M	0 20	06-E	P612	61	2006	0403		
OTHER	SOUR	CE(S):			CAS	REAC	T 14	5:41	9178								

OTHER SOURCE(S): CASREACT 145:419178

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$$R-N$$
 $N-Q$

AB The title compds. I [R = H, (un)substituted alkyl; Q = (un)substituted bicyclic aryl], useful as monoamine neurotransmitter re-uptake inhibitors, were prepared E.g., a multi-step synthesis of 2-(8-methyl-3,8-diazabicyclo[3.2.1]oct-3-yl)-6-nitroquinoline (II), starting from di-Et meso-2,5-dibromoadipate, was given. II showed IC50 of 16 μM , 4.6 μM and 0.0031 μM when tested for their ability to inhibit the reuptake of the monoamine neurotransmitters: dopamine, noradrenaline and serotonin in synaptosomes, resp. In other aspects the invention relates to the use of compds. I in a method for therapy and to pharmaceutical compns. comprising the compds. I.

MSTR 1

G2 = quinolinyl (opt. substd. by 1 or more G4)

G4 = 451 / 16

C(O)-G10 G5—C(O)—G6

G5 = NH G10 = NH2

Patent location: claim 1

Note: or pharmaceutically acceptable salts

Stereochemistry: and isomers and mixtures

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 26 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 145:397513 MARPAT

TITLE: Preparation of tetrahydroindazoles and analogs as

inhibitors of DNA gyrase and topoisomerase IV for the

treatment of bacterial infection

INVENTOR(S): Allison, Brett D.; Gomez, Laurent; Grice, Cheryl A.;

Hack, Michael D.; Morrow, Brian J.; Motley, Timothy
S.; Santillan, Alejandro; Shaw, Karen J.; Schwarz,
Kimberly L.; Tang, Liu Y.; Venkatesan, Hariharan;

Wiener, John J. M.

PATENT ASSIGNEE(S): Janssen Pharmaceutica N.V., Belg.

SOURCE: PCT Int. Appl., 172pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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WO 2006-US11631 20060330
     WO 2006105289
                       Α1
                            20061005
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
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     AU 2006230364
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                                            US 2006-393558
                       Α1
                                                             20060330
     EP 1863483
                                            EP 2006-748931
                       Α1
                            20071212
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     MX 200712234
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                            20080318
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                                           CN 2006-80019054 20071129
PRIORITY APPLN. INFO.:
                                            US 2005-667198P
                                                            20050331
                                            WO 2006-US11631
                                                            20060330
GΙ
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AB Bicyclic pyrazole compds. I [wherein B1, B5, B8 = (un)substituted CH or N; R2, R2, R6, R7 = H, (un)substituted alkyl, cycloalkyl, etc.; m = 0-1; n = 1-2; X = CH or N; Y = C(0), CH2C(0) or (un)substituted alkylene, etc.; A = (un)substituted (hetero)aryl; N1 or N2 is the anchoring site, with limitations] and isomers, racemates, tautomers, hydrates, solvates, pharmaceutically acceptable salts, esters, or amides thereof were prepared as antibacterial agents. For instance, tetrahydroindazole II was

synthesized in 30% yield by EDC/HOBt-mediated amidation of the corresponding benzodioxinecarboxylic acid with indazolamine in DMF. I showed inhibition against E. coli DNA gyrase and topoisomerase IV and antibacterial activity against both susceptible and resistant bacterial strains. Therefore, the invented compds. are useful for the treatment, prevention or inhibition of bacterial infection.

MSTR 1

G2 = 47 / N

_Ç----G3

G3 = 19 / 42 / CONH2

G4—G5 ₄C(0)—G11—G5

G4 = NH

G5 = cycloalkyl <containing 3-6 C> (opt. substd.)

G11 = NH

Patent location: claim 1

Note: substitution is restricted

Note: or tautomers, hydrates, solvates, pharmaceutically

acceptable salts, esters, or amides

Stereochemistry: or isomers or racemates

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 27 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 145:397380 MARPAT

TITLE: Preparation of 3,4-disubstituted coumarins and

quinolones for treatment of hepatitis C virus (HCV)

infection.

INVENTOR(S): Xu, Bin; Zhu, Qiang; Cho, Hyun-Joon; Fathi, Reza;

Yang, Zhen; Sandrasagra, Anthony; Liu, Yixin

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 74pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20060223783	A1	20061005	US 2005-93846	20050329
PRIORITY APPLN. INFO.	:		US 2005-93846	20050329
GI				

AB Title compds. (I; R1 = alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aralkyl, aryl, heteroaryl, halo, phosphate, phosphonate, etc.; 2 adjacent R1 may form a 5-6 membered (substituted) ring; n = 0-4; R2 = aralkyl, aryl, heteroaryl, cycloalkyl, cycloalkenyl; R3 = alkyl, cycloalkyl, alkenyl, alkynyl, aralkyl, aryl, heteroaryl, etc.; X = 0, NR4; Y = 0, NR5; R4, R5 = H, alkyl, cycloalkyl, alkenyl, alkynyl, aralkyl, aryl, heteroaryl, etc.), were prepared Thus, 4-hydroxycoumarin, 4-bromomethyltoluene, and K2CO3 were refluxed together in acetone overnight to give 10% I (X, Y = 0; R2, R3 = 4-MeC6H4CH2; n = 0). In an HCV replicon luciferase assay, the latter showed an IC50 = 8.29 $\mu \rm M$.

MSTR 1

G1 = 15

G2 = SO2

G3 = heteroaryl <containing zero or more N,

zero or more O, zero or more S> (opt. substd.)

G12 = 121

G13 = 83

83 (O)-G16

G16 = NH2 G21 = NH G23 = 103

N——G24

Patent location: claim 1

Note: or pharmaceutically acceptable salts or hydrates

L5 ANSWER 28 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 145:188748 MARPAT

TITLE: Preparation of quinolinium salts as anticancer drugs. INVENTOR(S): Macdonald, James E.; Hysell, Michelle K.; Yu, Dehua;

Li, Henry; Wong-Staal, Flossie

PATENT ASSIGNEE(S): Immusol Incorporated, USA SOURCE: PCT Int. Appl., 86 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	TENT	NO.		KII	.VD	DATE			Al	PPLI(CATI	и ис	ο.	DATE			
WO	2006	0787	54	A	1	2006	0727		M	O 20	06-U	S179.	3	2006	0118		
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		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KN,	KP,	KR,
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		MZ,	NA,	NG,	NΙ,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
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		CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	${ m ML}$,	MR,	NE,	SN,	TD,	ΤG,	BW,	GH,
		GM,	KE,	LS,	MW,	${ m MZ}$,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,	BY,
		KG,	KΖ,	MD,	RU,	ТJ,	TM										
AU	2006	2065	55	A.	1	2006	0727		Αl	J 20	06-2	0655	5	2006	0118		
CA	2595	224		A.	1	2006	0727		C	A 20	06-2	5952.	24	2006	0118		
EP	1841				_	2007							_	2006			
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			,	,										SI,		TR	
	1011			Α										2006			
	2008					2008								2006			
	2007													2007			
	2007					2007	1116			_	-			2007			
IORIT:	Y APP	LN.	INFO	.:					U	S 20	05-6	4509.	3P	2005	0118		

US 2005-715257P 20050908 WO 2006-US1793 20060118

GΙ

AB Title compds. e.g. [I; A = (substituted) Ph, heteroaryl; R = H, (substituted) alkyl, Ph, phenylalkyl; R1, R2 = H, CHO, cyano, (substituted) alkyl, (bicyclic) heterocyclyl, etc.], were prepared Thus, pyrvinium pamoate in CHCl3/EtOH at 50° was treated with H3PO4 in EtOH to precipitate pyrvinium phosphate. The latter showed IC50 <0.03 μM against MCF7 breast cancer cells in soft agar culture.

MSTR 2

G1 = 172

N 172 H⁺

G3 = CONH2G4 = CONH2 / 323

HN-G7

G7 = alkyl <containing 1-12 C> (opt. substd.)

Patent location: claim 25

Note: or pharmaceutically acceptable salts

Note: substitution is restricted

Note: additional ring formation also claimed

Stereochemistry: 90-E

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 29 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 145:145748 MARPAT

TITLE: Piperazinyl and piperidinyl ureas as modulators of

fatty acid amide hydrolase

INVENTOR(S): Apodaca, Richard; Breitenbucher, J. Guy; Pattabiraman,

Kanaka; Seierstad, Mark; Xiao, Wei Janssen Pharmaceutica N.V., Belg.

PATENT ASSIGNEE(S): Janssen Pharmaceutica N SOURCE: PCT Int. Appl., 151 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE			A:	PPLI	CATI	ON NO	ο.	DATE			
WO 2006074025	A 1	20060	0713		W	0 20	0.5-U	S473:	29	2005	1229		
W: AE, AG,									_			CA,	CH,
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VN, YU,	ZA, ZM	I, ZW											
RW: AT, BE,	BG, CH	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,
IS, IT,	LT, LU	, LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
CF, CG,	CI, CM	I, GA,	GN,	GQ,	GW,	$\mathrm{ML}_{m{\prime}}$	MR,	ΝE,	SN,	TD,	ΤG,	BW,	GH,
· · · · · · · · · · · · · · · · · · ·	LS, MW		•	SD,	SL,	SZ,	${\sf TZ}$,	UG,	ZM,	ZW,	AM,	AΖ,	BY,
, ,	MD, RU	,											
AU 2005322920										2005			
CA 2596393			-		_						-		
US 20060173184													
EP 1836179									_				
R: AT, BE,	•		•										•
·	LI, LI		LV,	MC,	ΝL,	PL,	P1,	RO,	SE,	SI,	SK,	IK,	AL,
ВА, НК, JP 2008526755	MK, YU		2724		т.	D 20	07 E	4960:	2	2005	1 2 2 0		
MX 200708134			-		-			4960. 134	-	2003			
IN 2007KN02653							-	134 N265:		2007			
NO 2007003923						_	-		_	2007			
KR 2007094635										2007			
CN 101146786		0000							-	2007			
ORITY APPLN. INFO		2000	0010							2004			
	. • •				_				-	2005			
						_ •			-				

AB Title compds. I [Z = N, CH; R1 = H, alkyl; Ar1 = (un)substituted 2-thiazolyl, 2-pyridinyl, 3-pyridinyl, 4-pyridinyl, 2-pyrimidinyl, 4-pyrimidinyl, 5-pyrimidinyl, Ph; Ar2 = (un)substituted 1-naphthyl, phenanthrenyl, pyrenyl, fluorenyl, 2-naphthyl, etc.; and their pharmaceutically acceptable salts, pharmaceutically acceptable prodrugs, and pharmaceutically active metabolites] were prepared as fatty acid amide hydrolase (FAAH) inhibitors. For example, reacting piperazine-1-carboxylic acid tert-Bu ester with Ph isocyanate, followed by Boc-deprotection and reductive alkylation with 2-naphthaldehyde gave piperazinyl urea II, which exhibited an IC50 of 17 nM in an FAAH assay. Thus, I and their pharmaceutical compns., are useful for treating disease states, disorders, and conditions mediated by FAAH, e.g., anxiety, pain, inflammation, sleep disorders, eating disorders, or movement disorders (such as multiple sclerosis).

MSTR 1B

$$G3 - G1 - C - R - G57$$

$$G3 - G1 - C - R - G57$$

$$G2 = 424 / 462$$

$$462 - 462$$

$$462 - 460$$

$$462 - 469$$

$$666 - 468$$

G6 = alkylamino <containing 1-4 C> / 54

G8---G9

G8 = C(0) / SO2

G9 = NH2G27 = 628

G8---G9 628

 $\mathsf{G35} \quad = 546-7 \ 547-460 \ 548-459 \ 594-458 \ 539-461$

G6 N 546 539 594 548

Patent location: claim 1

Note: or pharmaceutically acceptable salts, prodrugs, or

metabolites

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 30 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 145:145559 MARPAT

TITLE: Heteroaromatic quinoline compounds as

phosphodiesterase inhibitors, their preparation, pharmaceutical compositions, and use in therapy Verhoest, Patrick Robert; Helal, Christopher John;

Hoover, Dennis Jay; Humphrey, John Michael

PATENT ASSIGNEE(S): Pfizer Products Inc., USA

SOURCE: PCT Int. Appl., 90 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

INVENTOR(S):

PATENT NO	Э.	KI	ND	DATE			A.	PPLI	CATI	N NC	Ο.	DATE			
							_								
WO 20060	72828	A	2	2006	0713		M	O 20	05-I	в393	7	2005	1222		
WO 20060	72828	A	3	2006	1109										
W: A	AE, AG	, AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
(CN, CO	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
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F	KZ, LC	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,
I.	MZ, NA	NG,	ΝI,	NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,
	SG, SK	, SL,	SM,	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,
7	VN, YU	ZA,	ZM,	ZW											
RW: A	AT, BE	, BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,

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IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
            CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH,
            GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
            KG, KZ, MD, RU, TJ, TM
                                          AU 2005-323794
    AU 2005323794
                     A1
                           20060713
                                                           20051222
    CA 2592986
                           20060713
                                          CA 2005-2592986 20051222
                      Α1
    EP 1841757
                      Α2
                           20071010
                                          EP 2005-824101
                                                         20051222
        R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
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            BA, HR, MK, YU
    CN 101098866
                      Α
                           20080102
                                          CN 2005-80046085 20051222
                           20080724
                                          JP 2007-549961
    JP 2008526825
                      Τ
                                                           20051222
    US 20060154931
                           20060713
                                          US 2006-326221
                                                           20060105
                     A1
    NL 1030863
                           20060710
                                          NL 2006-1030863 20060106
                      Α1
    NL 1030863
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                          20061228
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                                          NO 2007-2918
                                                           20070607
                      Α
    IN 2007DN04794
                     Α
                          20070817
                                          IN 2007-DN4794
                                                           20070621
    KR 2007091005
                          20070906
                                          KR 2007-715460
                                                           20070705
                      Α
                         20070907
    MX 200708287
                      Α
                                          MX 2007-8287
                                                           20070706
                                          US 2005-642058P 20050107
PRIORITY APPLN. INFO.:
                                          WO 2005-IB3937
                                                           20051222
GΙ
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention relates to heteroaryl quinoline derivs. of formula I, which AB are phosphodiesterase (PDE) inhibitors, in some cases selective PDE-10 inhibitors. In compds. I, each R1 is independently selected from H, halo, OH, cyano, C1-8 alkyl, C2-8 alkenyl, C1-8 alkoxy, 4- to 7-membered heterocyclyl, etc.; p is 0-3; Het1 is (un)substituted mono- or bicyclic heteroaryl; Het2 is (un) substituted mono- or bicyclic heteroaryl, where Het2 is vicinal to the Ph ring on Het1; X1 and X2 are independently selected from O, S, (un)substituted N, and (un)substituted C, where are least one of X1 and X2 is C; and each Y is independently selected from N and (un) substituted C; provided that Het2 is not a tetrazole. The invention also relates to the preparation of I, pharmaceutical compns. comprising a compound I, as well as to the use of the compns. for the treatment of neurodegenerative and psychiatric disorders, such as psychosis. Substitution of 2-(chloromethyl)quinoline with Me 4-hydroxybenzoate followed by hydrolysis and amidation gave Weinreb amide II, which underwent addition of deprotonated 4-methylpyridine to give ketone III. Condensation of III with N-(dimethoxymethyl)-dimethylamine and heterocyclization with hydrazine gave pyrazole IV. The compds. of the invention express IC50 values for PDE-10 inhibition of less than 10 μM (no specific data).

MSTR 1

10/572,914

G24 = 64

G25 = alkylamino <containing 1-8 C> / 56

C(0)-G31

G31 = NH2

Patent location: claim 1

Note: substitution is restricted

L5 ANSWER 31 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 144:488666 MARPAT

TITLE: Preparation of quinoline, tetrahydroquinazoline, and

pyrimidine derivatives as MCH antagonist for treatment

of CNS disorders

INVENTOR(S): Sekiguchi, Yoshinori; Kanuma, Yukihiro; Omodera,

Katsunori; Busujima, Takeshi; Tran, Thuy-Ahn; Han, Sangdong; Casper, Martin; Brian, A. Kramer; Semple,

Graeme; Zou, Ning

PATENT ASSIGNEE(S): Taisho Pharmaceutical Co., Ltd., Japan; Arena

Pharmaceutical Inc.

SOURCE: Jpn. Kokai Tokkyo Koho, 781 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 2006124387 A 20060518 JP 2005-286311 20050930
PRIORITY APPLN. INFO.: JP 2004-287659 20040930
GI

AB Title compds. [I, II, III; wherein R1 = (un)substituted (cyclo)alkyl, (cyclo)alkenyl, alkynyl, aryl; R2 = H, halo, OH, carboxy, carbamoyl, amino, (un)substituted alkyl, alkoxy; T = independently H, halo, OH, carboxy, carbamoyl, amino, cyano, NO2, alkenyl, alkynyl, cycloalkyl, (un)substituted alkyl, alkoxy; p = 0-5; L = aminocycloalkylideneamino, etc.; Y = bond, CH2, CO2, OCO, SO2, CO, CS, CONH, CSNH, etc.; with provisos; and pharmaceutically acceptable salts, hydrates, or solvates thereof] were prepared as antagonists of melanin concentrating hormone (MCH),

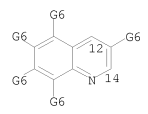
an

endogenous ligand of G-protein coupled receptors (GPCRs). Examples include solution and solid phase general synthetic methods and phys. data for nearly 3400 invention compds. In addition, all exemplified compds. were assayed using high throughput functional screening to detect intracellular Ca2+ concns. for accessing GPCR activation. For instance, reaction of 2,4-dichloro-6-methylpyrimidine with dimethylamine gave 2-chloro-4-(dimethylamino)-6-methylpyrimidine (40%), which was coupled with cis-(4-aminocyclohexyl)carbamic acid tert-Bu ester (60%). Deprotection (72%), amidation, and workup provided the benzamide (IV) •TFA. The latter demonstrated MCH antagonist activity with an IC50 value of 7.6 nM. Thus, pharmaceutical compns. comprising I are useful for the prophylaxis or treatment of improving memory function, sleeping and arousal, anxiety, depression, mood disorders, seizure, obesity, diabetes, appetite and eating disorders, cardiovascular disease, hypertension, dyslipidemia, myocardial infarction, binge eating disorders including bulimia, anorexia, mental disorders including manic depression, schizophrenia, delirium, dementia, stress, cognitive disorders, attention deficit disorder, substance abuse disorders, and dyskinesias including Parkinson's disease, epilepsy, and addiction (no data).

10/572,914

MSTR 1

G1 = 12-5 14-2



G2 = NHNH2 G6 = CONH2

Patent location: claim 1

Note: substitution is restricted

Note: additional substitution also claimed

L5 ANSWER 32 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 144:360024 MARPAT

TITLE: Colored hardenable composition for color filter and

production method of color filter

INVENTOR(S): Kato, Yasuhiro; Seto, Nobuo; Mizukawa, Hiroki;

Fujimori, Toru

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 72 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 2006091190 A 20060406 JP 2004-274216 20040921

PRIORITY APPLN. INFO:: JP 2004-274216 20040921

GI

$$AN = N - N - R^{205}$$

$$R^{206}$$

AB The invention relates to a colored hardenable composition, suited for use in making a color filter of a solid state camera and a liquid crystal display, comprising compds. represented by I [A = five member heterocyclic residue; B1 = CR201 or N, B2 = CR202 or N, and B1 and B2 may not be N simultaneously; R205 and R 206 = H, aliphatic, aromatic, etc., and R205 and R206

may not be H simultaneously; G, R201, and R202 = H, halo, aliphatic, aromatic, etc.; R202 and R205, and/or R205 and R206 may join to form a 5 or 6 member ring] and II [R303, R304, R307 and R308 = H, halo, aliphatic, aromatic, etc.; R301, R302, R305, and R306 = C, H, halo, aliphatic, etc., and R301, R302 and R305, R306 may form a 5- or 6-member carbon ring; m and n = 0-4 integer].

MSTR 2

G5 = CONH2 (opt. substd.) / acylamino / SO2NH2 (opt. substd.)

G1 + G4 = CH = CHCH = CH (opt. substd. by 1 or more G5)

Patent location: claim 1

Note: additional ring formation also claimed

L5 ANSWER 33 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 144:350543 MARPAT

TITLE: Preparation of indole derivatives as inhibitors of

interaction between MDM2 and p53

INVENTOR(S): Lacrampe, Jean Fernand Armand; Meyer, Christophe;

Ligny, Yannick Aime Eddy; Csoka, Imre Christian Francis; Van Hijfte, Luc; Arts, Janine; Schoentjes, Bruno; Wermuth, Camille Georges; Giethlen, Bruno;

Contreras, Jean-Marie; Joubert, Muriel Janssen Pharmaceutica N.V., Belg.

PATENT ASSIGNEE(S): Janssen Pharmaceutica N.V., Belg SOURCE: PCT Int. Appl., 132 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	CENT :				ND	DATE								DATE			
	WO	2006				1									2005	0916		
		W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KP,	KR,	KΖ,
			LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,
			NA,	NG,	NI,	NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,
			SK,	SL,	SM,	SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,
			YU,	ZA,	ZM,	ZW												
		RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	IE,
			IS,	ΙΤ,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
			CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG,	BW,	GH,
			GM,	ΚE,	LS,	MW,	MZ,	NΑ,	SD,	SL,	SZ,	ΤZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,	BY,
							ТJ,											
	AU 2005286525										-			_	2005			
		2579							_									
	ΕP	1809	-												2005			
		R:													GB,			
			,	,	,	•	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	AL,
			,	,	MK,													
		1010		_			2007			_					2005			
		2008					2008					07-5			2005			
		2005		-			2008					05-1			2005			
		2008			А		2008					07-5			2007			
		2007		A		2007								2007				
	MX 200703375						2007								2007			
	KR 2007058622 DRITY APPLN. INFO						2007	0608							2007			
PRIOF	RITY	APP	LN.	INFO	.:							04 - 7			2004			
															2004			
										W	0 20	05-E	P546	0 4	2005	0916		
GI																		

The title compds. I [wherein m = 0-2; n = 0-3; p, q and q' = independently 0 or 1; X = CO or (un)substituted CH2; Q-Y = (un)substituted CH=C, CO-CH, CO-N, CH2-CH, or CH2-N; R1 = H, aryl, heteroaryl, alkyl, etc.; R2 = H, halo, alkyl, alkoxy, etc.; R3 = H, alkyl, heteroaryl, etc.; R4 and R5 = independently H, halo, alkyl, CN, etc.; R6 = H, alkoxycarbonyl, or alkyl; Z = (un)substituted heteroaryl; with provisos] or N-oxides, salts, or stereoisomers thereof are prepared as inhibitors of interaction between MDM2 and p53. For example, the compound II●xHCl was prepared in a multi-step synthesis. I showed inhibitory effect on cell proliferation. Formulations containing I as an active ingredient were also described.

MSTR 1

G15—G27

G15 = 13

G16 = phenylene (opt. substd. by (up to 1) G17) G18 = NH

G27 = 180

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G28
G28
G28
N G28
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G28 = 298 / CONH2 (opt. substd.)

C(0)-G29

G29 = heteroaryl <containing up to 14 atoms,

1-5 heteroatoms, zero or more N, zero or more O, zero or more S (no other heteroatoms), 1-3 rings>

(opt. substd.)

Patent location: claim 1

Note: and N-oxides or addition salts

Note: additional ring formation also claimed

Note: also incorporates claim 10 Stereochemistry: or sterochemical isomers

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 34 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 144:232928 MARPAT

TITLE: Preparation of heterocyclic compounds as novel

antimalaria agents

INVENTOR(S): Nakamoto, Kazutaka; Matsukura, Masayuki; Tanaka,

Keigo; Inoue, Satoshi; Tsukada, Itaru; Haneda, Toru;

Ueda, Norihiro; Abe, Shinya; Sagane, Koji

PATENT ASSIGNEE(S): Eisai Co., Ltd., Japan SOURCE: PCT Int. Appl., 326 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2006016548 A1 20060216 WO 2005-JP14505 20050808

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW

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KG, KZ, MD, RU, TJ, TM
     WO 2005033079
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             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
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     EP 1782811
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                                         EP 2005-768893 20050808
                     A1
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             BA, HR, MK, YU
     IN 2007DN00839
                    Α
                           20070803
                                           IN 2007-DN839
                                                            20070131
PRIORITY APPLN. INFO.:
                                           JP 2004-232617
                                                            20040809
                                           WO 2004-JP14063 20040927
                                          JP 2005-82760
                                                            20050322
                                           JP 2003-342273
                                                            20030930
                                          JP 2004-68186
                                                            20040310
                                          WO 2005-JP14505 20050808
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GΙ

$$A^1$$
 X^1 CH_2 E

AB Antimalaria agents containing compds. represented by the formula (I) (wherein A1 = each optionally substituted 3-pyridyl or 6-quinolyl; X1 = $-C(:Y1)-NH-; Y1 = 0; E = each optionally substituted furyl, thienyl, or phenyl; provided that A1 may have one to three substituents and E has one or two substituents), salts of the compds., or hydrates of either are disclosed. Thus, a solution of 2-aminonicotinic acid and [[5-(3-chlorobenzyl)furan-2-yl]methyl]amine in DMF was treated with benzotriazol-1-yl-tris(dimethylamino)phosphonium hexafluorophosphate and Et3N and stirred at 80° for 40 min to give 2-amino-N-[5-(3-chlorobenzyl)furan-2-ylmethyl]nicotinamide (II). II showed min. inhibitory concentration of 6.25 <math>\mu$ g/mL against yeast expressing plasmodium GWT1 gene (opfGWT1).

MSTR 1

$$G1 \longrightarrow C \longrightarrow NH \longrightarrow CH_2 - G2$$

$$G1 = 284$$

= CONH2 / alkylamino <containing 1-6 C>

(opt. substd.)

Patent location: claim 1

Note: or salts or hydrates

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 4 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 35 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 144:150653 MARPAT

Preparation of dipeptide analogs as hepatitis C TITLE:

inhibitors

Bailey, Murray, D.; Bhardwaj, Punit; Ghiro, Elise; INVENTOR(S):

Goudreau, Nathalie; Halmos, Teddy; Llinas-Brunet, Montse; Poupart, Marc-Andre; Rancourt, Jean

PATENT ASSIGNEE(S): Boehringer Ingelheim International GmbH, Germany;

Boehringer Ingelheim Pharma GmbH & Co. KG

SOURCE: PCT Int. Appl., 153 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

E	PATENT	NO.		KI	ND	DATE			A.		CATI		٥.	DATE			
_ ⊽	WO 200	060077	00	 A	1	2006	0126		M				 5	2005	0715		
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KM,	KP,	KR,	KΖ,
		LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,
		NG,	NΙ,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,
		SL,	SM,	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,
		ZA,	ZM,	ZW													
	RV	∛: AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	IE,
		IS,	ΙΤ,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG,	BW,	GH,
		GM,	KE,	LS,	MW,	${ m MZ}$,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,	BY,
		KG,	KΖ,	MD,	RU,	ТJ,	TM										
	CA 25	73219		A	1	2006	0126		C.	A 20	05-2	5732	19	2005	0715		
E	EP 17	71453		А	1	2007	0411		E:	P 20	05-7	6353	9	2005	0715		
	R:	AT,	BE,	ВG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	IE,
							•		•					SI,		TR	
	JP 200																
J	JS 200	060046	965	А	1	2006	0302		U	S 20	05-1	8567	1	2005	0719		
PRIORI	ITY AE								2004								
									M	0 20	05-C.	A111	5	2005	0715		

OTHER SOURCE(S): CASREACT 144:150653

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention relates to peptides I [m, n are 1 or 2; R1 is (halo)alkyl, (halo)alkenyl or (halo)alkynyl; R2 is NH-R5, O-R5, S-R5, SOm-R5, OCH2-R5 or CH2O-R5, where R5 is (un)substituted aryl or heterocyclyl; R3 is carboxylic ester, carbamoyl, sulfinyl, sulfonyl or acyl groups; R4 is (un)substituted alkyl, alkenyl, cycloalkyl, aryl or heterocyclyl (with provisos)] (or racemates, diastereomers or salts) for the treatment of hepatitis C viral infection. Thus, dipeptide II was prepared via peptide coupling reactions in solution and etherification of a hydroxyproline intermediate. Many peptides I have IC50 values < 0.5 μM in the NS3-NS4A protease assay and < 1 μM in the cell-based luciferase HCV RNA replication assay.

MSTR 1

$$G2 = NH$$

 $G20 = 88$

$$G21 = CONH2$$

 $G30 = 89$

Patent location: claim 1

Note: additional substitution also claimed

Note: or salts

Stereochemistry: or racemates, diastereomers, and optical isomers

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 36 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 144:45455 MARPAT

TITLE: Tricyclic compounds as inhibitors of the hypoxic

signaling pathway for cancer treatment

INVENTOR(S): Melillo, Giovanni; Shoemaker, Robert H.; Cardellina,

John H.; Currens, Michael J.; Creighton-Gutteridge, Mark; Uranchimeg, Badarch; Rapisarda, Annamaria;

Scudiero, Dominic A.

PATENT ASSIGNEE(S): The Government of the United States of America as

Represented by the Secretary, Department of Health,

USA

SOURCE: PCT Int. Appl., 119 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	PATENT NO.				ND	DATE			APPLICATION NO.					DATE			
				A2 A3		20051215		WO 2005-US16569 20050511									
WO		AE,	AG,	AL,	AM,	AT, CZ,	AU,	•	•	•	•	•	•	•	•	•	•
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KP,	KR,	KZ,
		,	,	,	,	LT, OM,	,	,	,	,	,	,	,	,	,	,	,
			SM, ZM,		TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,
	RW:	BW,	GH,	GM,		LS,	•	•	•	•	•	•	•	,	•	•	•
		•	•	•	•	MD, GB,	•	•	•	•	•	•	•	•	•	•	•
		,	,	,	,	TR, TG	BF,	BJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,
D T TT	7 7 7 7	MR, NE, SN, TD, TG															

PRIORITY APPLN. INFO.:

US 2004-570615P 20040512 US 2004-618279P 20041012

GΙ

AB Tricyclic compds. I (wherein X and Y are independently O, S, N, NR4, CR5 or CR6R7; R1 = one or more substituents independently selected from acyl, acyloxy, alkoxy, alkyl, alkylthio, amino, aryl, aza, CO, carboxamide, diamine, halogen, OH, mercapto, NO, sulfonyl, sulfonamido and sulfato, at least one of which is carboxamide or diamine; R2 and R3 are either joined

to form an (un)substituted six-membered aromatic ring, or one of R2 and R3 is an (un)substituted aryl group; R4, R5, R6 and R7 are independently H or a substituent as defined for R1 above) or II (wherein R8 is defined the same as R1 above; Ar = an (un)substituted aryl group; W and Z = NR9 or =N-; and R9 = H or a substituent as defined for R1 above) that selectively inhibit HIF-1 α activity are disclosed. Methods also are disclosed for reducing HIF-1 α activity, and for inhibiting angiogenesis, tumorigenesis and/or metastasis, in a subject. In some embodiments, the tricyclic compds. surprisingly inhibit HIF-1 α activity at non-cytotoxic concns., thereby avoiding drug side effects associated with significant cytotoxicity.

MSTR 1

$$G1 = 20 / 26 / 41$$

G6 = any ring <containing 5 or more atoms,
 zero or more N, zero or more O,
 zero or more S (no other heteroatoms), aromatic,
 2 or more double bonds>

G7 = NH2 / 28

$$G9 = 37$$

$$G12 = 39$$

$$G13 = 56$$

 $G14 = 182-1 \ 183-4 \ 185-57 \ 186-69$

G18 = NH / SO2 G19 = 42

G11-G12

Patent location: claim 1

Note: all ring carbons can also be nitrogen

Note: substitution is restricted

L5 ANSWER 37 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:347191 MARPAT

Preparation of benzyl pyridazinone derivatives as TITLE:

non-nucleoside reverse transcriptase inhibitors

INVENTOR(S): Dunn, James Patrick; Elworthy, Todd Richard; Hogg,

Joan Heather; Stefanidis, Dimitrios F. Hoffmann-La Roche A.-G., Switz.

PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 58 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT I			KI	ND	DATE			A.	PPLI	CATI	ON N	ο.	DATE				
WO 2005	0903:	17	А	1	2005	0929		W	0 20	05-E	P277	9	2005	0316			
W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	KP,	KR,	KΖ,	LC,	
	LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NΙ,	
	NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	
	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
RW:	BW,	GH,	GM,	KΕ,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	
	ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,	IS,	ΙΤ,	LT,	LU,	MC,	NL,	PL,	PT,	
	RO,	SE,	SI,	SK,	TR,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	${ m ML}$,	

MR, NE, SN, TD, TG 20050929 CA 2005-2559552 20050316 CA 2559552 A1 EP 1730120 20061213 EP 2005-716102 20050316 Α1 R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR CN 1934092 20070321 CN 2005-80008974 20050316 Α JP 2007530477 Τ 20071101 JP 2007-504308 20050316 US 20050215554 Α1 20050929 US 2005-85869 20050322 US 7288542 В2 20071030 PRIORITY APPLN. INFO.: US 2004-555798P 20040323 WO 2005-EP2779 20050316 OTHER SOURCE(S): CASREACT 143:347191

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AΒ Title compds. I [R1, R2, R3 and R4 independently = H, alkyl, haloalkyl, etc.; R5 = (un)substituted aryl or heteroaryl; R6 = (CH2)pOH, CH2CO2R9, CH2OP(O)(OH)2, etc.; R7 and R8 independently = H, amino, alkylamino, etc.; R9 = H or alkyl; p = 1-3] and their pharmaceutically acceptable salts, are prepared and disclosed as non-nucleoside reverse transcriptase (nnRT) inhibitors. Thus, e.g., II was prepared by alkylation of III with formaldehyde. The pharmacokinetic activity was evaluated by orally administering various doses of I to Hanover-Wistar rats and subsequent determination of test compound concentration using HPLC and it was revealed that selected

compds. of the invention possessed Cmax values in the range of 2.2 up to 15.5 μ g/mL. I as non-nucleoside reverse transcriptase inhibitors should prove useful in the treatment of HIV mediated diseases. Pharmaceutical compns. comprising I are disclosed.

MSTR 1

$$G5 = 193$$

G11 = alkylamino <containing 1-6 C> / CONH2

10/572,914

G14 = N

Patent location: claim 1

Note: and pharmaceutically acceptable acid or base

addition salts, hydrates, solvates, or clathrates

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ΙI

L5 ANSWER 38 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:336409 MARPAT

TITLE: Dye-containing photosensitive material compositions

for color filters in solid-state image pickup and in

liquid crystal displays

INVENTOR(S): Kato, Yasuhiro; Mizukawa, Hiroki PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 60 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 2005258093 A 20050922 JP 2004-69741 20040311

PRIORITY APPLN. INFO.: JP 2004-69741 20040311

GI

$$A-N=N$$

$$R^{1}=B^{2}$$

$$N$$

$$R^{5}$$

$$R^{6}$$

AB The title composition contains magenta dye I(R1-2 = H, substituent; m = integer 0-2; n, j = integer 0-4; Y = 0, N, C; Z = C, N, O, S) and yellow dye <math>II(A = 5-membered heterocyclic ring; B1-2 = -CR7=; -CR8=, N; R5-6 = H, aliphatics, aroms., etc.; G = H, halo, aliphatics, aroms., etc.). The composition shows good storageability and provides red color of light- and heat-resistance.

MSTR 1

G1 = acylamino / CONH2 / SO2NH2 Patent location: claim 1

L5 ANSWER 39 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:266946 MARPAT

TITLE: Preparation of pyridines and related compounds as

TGF-eta inhibitors

INVENTOR(S): Shimizu, Kiyoshi; Shimizu, Toshiyuki; Kawakami,

Kazuki; Nakoji, Masayoshi; Sakai, Teruyuki

PATENT ASSIGNEE(S): Kirin Beer Kabushiki Kaisha, Japan

SOURCE: PCT Int. Appl., 461 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.				KI:	ND :	DATE			A.	PPLI	CATI	N NC	0.	DATE			
	WO	2005	0803	 77	A	1	2005	0901		W	0 20	05-J:	P261	0	2005	0218		
		W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,
			LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	ΝA,	NI,
			NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
			ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
		RW:	BW,	GH,	GM,	ΚE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,
			AΖ,	BY,	KG,	KΖ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
			EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IS,	IT,	LT,	LU,	MC,	NL,	PL,	PT,
			RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,
			MR,	ΝE,	SN,	TD,	ΤG											
	ΕP	1724	268		Α	1	2006	1122		E	P 20	05-7	1928	0	2005	0218		
		R:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,
			IS,	ΙΤ,	LI,	LT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR		
PRIOF	ORITY APPLN. INF			INFO	.:					J.	P 20	04 - 4	5383		2004	0220		
										W	0 20	05-J	P261	0	2005	0218		
GI																		

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Title compds. I [A = II; Z = O, etc.; D1, D2, D3, D4, X, E, G, J, L, M = C, N; further details on D1, D2, D3, D4, X, E, G, J, L, M are given.; R1-R6, R10-R14 = H, halo, etc.] were prepared For example, reaction of

10/572,914

4-chloro-6,7-dimethoxyquinazoline with 5,6-dimethyl-[2,2'-bipyridin]-3-ol, e.g., prepared from 2,3-dimethylfuran in 2 steps, afforded compound III in 81% yield. In TGF- β signal inhibition assays (in vitro), compound III exhibited the inhibitory activity of 89% at 1 μ M. Compds. I are claimed useful for the treatment of arthritis, ulcer, etc.

MSTR 1

G1 = 19

G2 = NH G4 = 32

C (O)-G5

G5 = NH2 / heterocycle <containing 3-9 atoms, 1 or more N, zero or more O, zero or more S (no other heteroatoms), attached through 1 or more N, non-aromatic, saturated> (opt. substd.)

G8 = 75

75——G4

G12 = 89

613 g

G13 = 139

1860)-G5

G16 = 7

Patent location: claim 1

Note: or pharmaceutically acceptable salts or solvates

Note: additional ring formation also claimed

Note: substitution is restricted also incorporates claim 68

REFERENCE COUNT: 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 40 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:248301 MARPAT

TITLE: Preparation of substituted quinolines as MTP/Apo-B

secretion inhibitors for treating obesity and

associated conditions

INVENTOR(S): Bertinato, Peter; Couturier, Michel Andre; Hamanaka,

Ernest Seiichi; Ewing, Marcus Douglas; Robinson, Ralph

Pelton, Jr.; Tickner, Derek Lawrence

PATENT ASSIGNEE(S): Pfizer Products Inc., USA SOURCE: PCT Int. Appl., 162 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PAT	TENT	NO.		KI	ND	DATE			A	PPLI	CATI	ON N	Ο.	DATE			
WO	2005	0803	73	A	1	2005	0901		W	20	05-I	 В167		2005	0124		
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	KP,	KR,	KΖ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NΙ,
		NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	${ m TZ}$,	UG,	ZM,	ZW,	AM,
		ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM,	ΑT,	BE,	ВG,	CH,	CY,	CZ,	DE,	DK,
					•		•							MC,			•
							BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,
			ΝE,														
_										-		_	-	2005	-		
-									_	-				2005	-		
EΡ														2005			
	R:								•		•			NL,	•		
		,	•	•		FΙ,	RO,	MK,	CY,	AL,	TR,	ВG,	CZ,	EE,	HU,	PL,	SK,
			HR,	,													
	1914								_					2005			
	2005													2005			
	2007								_	_				2005	-		
US	2005	0234	099	A	1	2005	1020		U	S 20	05 - 4	9852		2005	0203		

NL	1028192	A1	20050808	NL	2005-1028192	20050204
NL	1028192	C2	20060530			
US	20060223851	A1	20061005	US	2006-424488	20060615
US	7368573	В2	20080506			
MX	2006PA07785	A	20060926	MX	2006-PA7785	20060706
IN	2006DN03919	A	20070427	ΙN	2006-DN3919	20060707
KR	799802	В1	20080131	KR	2006-715770	20060803
NO	2006003928	A	20061031	NO	2006-3928	20060901
US	20070093525	A1	20070426	US	2006-554351	20061030
US	7393958	В2	20080701			
PRIORIT	Y APPLN. INFO.:			US	2004-541678P	20040204
				US	2004-633763P	20041206
				WO	2005-IB167	20050124
				US	2005-49852	20050203

$$(R^{7})_{n}$$

$$(R^{2})_{m}$$

$$R^{3}$$

$$R^{4}$$

$$R^{6}$$

$$R^{5}$$

$$R^{5}$$

AΒ This invention relates to MTP/Apo-B secretion inhibitors of Formula (I) wherein R1-R7, X1, m and n are as defined below, as well as pharmaceutical compns. comprising the compds., and methods of use of the compds. and compns. The compds. of the invention are useful in treating obesity and associated diseases, conditions or disorders. For I the variables are: R1 = substituted Ph or pyridine; m = 0-2; n = 0-4; X1 = N or C(Rb) where Rb = Hor R7; R2, R7, and R9 = halo, OH, CN, alkyl, alkoxy, alkoxyalkyl, halo-substituted alkyl, halo-substituted alkoxy, alkylthiobenzyloxy, hydroxyalkyl, alkenyl, alkynyl, C(0)N(Rc)(R11), N(R11)C(0)R12, N(R11)CO2R12, N(R11)S(O)SR12, C(O)R12, CO2R12, OC(O)R12, SO2N(Rc)(R11) and S(0)vR12; Rc = H or alkyl; s = 1-2; v = 0-2; R3 and R4 = H or taken together with the C to which they are attached form a carbonyl group; R5 and R10 = H, alkyl, halo-substituted alkyl, cycloalkyl, C(0)R12, alkoxyalkyl, alkylthioalkyl and SO2R12.;. Variables for I continued: R6 = optionally subsituted alkyl, pyridyl, Ph, phenylalkyl, alkenyl, alkynyl, CH2N(Rc)(R13), C(0)N(R14)(R15), CO2R2O or CH2-W-Y where W=O or S; and Y = H, alkyl, cycloalkyl, optionally substituted cycloalkylalkyl, Ph and phenylalkyl; R11 = H, alkyl, halo-substituted alkyl, cycloalkyl, alkoxyalkyl and alkylthioalkyl; R12 = optionally substituted alkyl or cycloalkyl, group; R13 = alkyl, phenylmethyl, C(0)R16 and S(0)2R16; R14 = H, optionally substituted alkyl, cycloalkyl, cycloalkylalkyl, Ph and phenylalkyl; R15 = H, optionally substituted alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkylalkyl, Ph, phenylalkyl, pyridyl, pyridylalkyl, C(0)R12 and SO2R12; or R15 = (CH2)tN(R17)(R18) where t = 2-4 and R17 and ${\tt R18}$ together with the N to which they are attached to form a heterocyclic ring, which is optionally substituted; or R14 and R15 together with the N to which they are attached to form a heterocyclic ring which is optionally

substituted; and R16 = optionally substituted alkyl, Ph or phenylalkyl.

MSTR 1

G2 = 46

G24-C(O)-R

G3 = C(O) G5 = NH

G21 = CONH2 (opt. substd.) G24 = NH (opt. substd.)

Patent location: claim 1

Note: or pharmaceutically acceptable salts, or prodrugs

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 41 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:211934 MARPAT

TITLE: Preparation of 4-heteroaryloxy-6-piperazinopyrimidines

as vanilloid receptor ligands

INVENTOR(S): Wang, Hui-ling; Balan, Chenera; Doherty, Elizabeth M.;

Falsey, James R.; Gore, Vijay Keshav; Katon, Jodie;

Norman, Mark H.

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 46 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 20050176726	A1	20050811	US 2005-56568	20050211
AU 2005212517	A1	20050825	AU 2005-212517	20050211
CA 2555685	A1	20050825	CA 2005-2555685	20050211
WO 2005077944	A1	20050825	WO 2005-US4378	20050211
W: AE, AG,	AL, AM	, AT, AU, AZ,	BA, BB, BG, BR, BW	, BY, BZ, CA, CH,

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CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
             EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT,
             RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,
             MR, NE,
                     SN, TD, TG
     EP 1720868
                      Α1
                           20061115
                                            EP 2005-722962
                                                             20050211
            AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA,
             HR, LV, MK, YU
                             20070425
                                            CN 2005-80008675 20050211
     CN 1953976
                       Α
     BR 2005007927
                             20070717
                                            BR 2005-7927
                                                             20050211
                       Α
     JP 2007522235
                             20070809
                                            JP 2006-553265
                       Τ
                                                             20050211
     MX 2006PA09059
                             20061019
                                            MX 2006-PA9059
                       Α
                                                              20060809
     KR 2007033325
                       Α
                             20070326
                                            KR 2006-718172
                                                              20060906
     KR 813093
                       В1
                             20080317
     NO 2006004055
                       Α
                             20061024
                                            NO 2006-4055
                                                              20060908
PRIORITY APPLN. INFO.:
                                            US 2004-543896P
                                                             20040211
                                            WO 2005-US4378
                                                             20050211
                         CASREACT 143:211934
OTHER SOURCE(S):
GΙ
```

AB The title compds. I [X = N, C; R1 = (un)substituted (un)saturated 5-7 membered ring containing 1-4 atoms selected from N, O and S; R2 = (un)substituted partially saturated or unsatd. 8-11 membered bicyclic ring containing 1-4 atoms selected from N, O and S; R31, R32 = H, Me, Et; or R31 and R32 together may be combined with the carbon atom to which they attached to form cyclopropyl; R4 = H, Me], useful for the treatment of acute, inflammatory and neuropathic pain, dental pain, general headache, migraine, cluster

headache, mixed-vascular and non-vascular syndromes, tension headache, general inflammation, arthritis, rheumatic diseases, osteoarthritis, inflammatory bowel disorders, inflammatory eye disorders, inflammatory or unstable bladder disorders, psoriasis, skin complaints with inflammatory components, chronic inflammatory conditions, inflammatory pain and associated hyperalgesia and allodynia, neuropathic pain and associated hyperalgesia and allodynia, diabetic neuropathy pain, causalgia, sympathetically maintained pain, deafferentation syndromes, asthma, epithelial tissue damage or dysfunction, herpes simplex, disturbances of visceral motility at respiratory, genitourinary, gastrointestinal or vascular regions, wounds, burns, allergic skin reactions, pruritus, vitiligo, general gastrointestinal disorders, etc., were prepared E.g., a multi-step synthesis of II, starting from 4,6-dichloropyrimidine and 2-aminobenzothiazol-4-ol, was given. Compds. I were tested to evaluate their properties at human VR1 (data given for representative compds. I). The pharmaceutical composition comprising the compound I is disclosed.

MSTR 1

$$G7 = 55$$

$$G11 = 59-52 60-50$$

$$G12 = 61$$

$$G13 = 532 / 538 / 556$$

$$G22 = 124 / SO2$$

____G23 124

G23 = O / NH

G24 = NH2G35 = 558

G32-G24

Patent location: claim 1

Note: or pharmaceutically acceptable salts or hydrates

Note: substitution is restricted

L5 ANSWER 42 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:172772 MARPAT

TITLE: Preparation of quinoline derivatives as MCH modulators

Evertsson, Emma; Inghardt, Tord; Lindberg, Jan; INVENTOR(S):

Linusson, Anna; Giordanetto, Fabrizio

Astrazeneca Ab, Swed. PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 114 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PA:	TENT				MD.	DATE			A.	PPLI	CATI	ON N	0.	DATE			
WO	2005				1	2005	0721		M	0 20	 05-s:	 Е4		2005	0105		
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:													UG,			
		AZ,	BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,	IS,	ΙT,	LT,	LU,	MC,	NL,	PL,	PT,
		RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,
		MR,	NE,	SN,	TD,	ΤG											
EP	1706	384		A	1	2006	1004		E.	P 20	05-7	0467	8	2005	0105		
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	FI,	RO,	CY,	TR,	BG,	CZ,	EE,	HU,	PL,	SK,	IS		
CN	1906	169		Α		2007	0131		C1	N 20	05-8	0001	921	2005	0105		
JP	2007	5178	68	T		2007	0705		J:	P 20	06-5	4918	4	2005	0105		
IN	2006	DN03	548	Α		2007	0817		I	N 20	06-D	N354	8	2006	0620		
US	2007	0185	079	А	1	2007	0809		U	S 20	06-5	9699	4	2006	1122		
ORIT	Y APP	LN.	INFO	.:					G:	В 20	04 - 1	96		2004	0107		
									G:	В 20	04-2	5209		2004	1116		
									M	O 20	05-S	E4		2005	0105		
ER SO	DURCE	(S):			CAS	REAC'	Т 14.	3:17.	2772								

$$(R^{2})_{m}$$
 R^{4}
 $N-L^{1}-N-L^{2}-R^{5}$
 R^{3}
 R^{3}

AB Title compds. I [R1 = (un)substituted alkoxy, alkyl, NRaRb, etc.; R2 = (un)substituted alkoxy, alkyl, NRaRb, etc.; Ra and Rb independently = H, alkyl or Ra and Rb together with the nitrogen to which they are attached from a 3-7 membered heterocycle optionally including 0; n = 0-3; m = 0-1; R3 = H or alkyl; L1 = (CH2)pcycloalkyl(CH2)q with provisions; p and q independently = 0-1; R4 = H or (un)substituted alkyl; L2 = (un)substituted (CH2)x or 5-6 membered carbocycle fused to R5; x = 1-3; R5 = (un)substituted Ph, naphthyl, heterocycle, etc.] and their pharmaceutically acceptable salts, are prepared and disclosed as melanin concentrating hormone (MCH) modulators. Thus, e.g., II was prepared by palladium

catalyzed coupling of benzyl[(1R,2S,4S,6S)-6-aminobicyclo[2.2.1]hept-2-yl]benzylcarbamate (preparation given) with 2-chloro-6-methoxy-4-methylquinoline followed by deprotection and subsequent reductive alkylation with thiophene-3-carbaldehyde. The activity of I was evaluated in MCH1 receptor radioligand binding assays and it was revealed that compds. of the invention displayed IC50 values of less than 2 μM . I as MCH modulator should prove useful in the treatment of obesity, anxiety and depression. Pharmaceutical compns. comprising I are disclosed.

MSTR 1

10/572,914

G1 = 15

C(O)-G2

G2 = NH2 / heterocycle < containing 3-7 atoms,

1 or more N, attached through 1 N, non-aromatic, saturated>

G3 = alkylamino <containing 1-4 $\stackrel{\frown}{C}>$ / 19

C(O)-G2

Patent location: claim 1

Note: substitution is restricted Note: also incorporates claim 17

Note: and pharmaceutically acceptable salts Stereochemistry: and optical isomers and racemates

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 43 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:123108 MARPAT

TITLE: Pyrazolylazoquinolines, their chelates, and WORM disks

with high-speed and -density recording

INVENTOR(S): Ueno, Yasunobu; Sato, Tsutomu; Tomura, Tatsuya;

Noguchi, Shu

PATENT ASSIGNEE(S): Ricoh Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 29 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 2005179418 A 20050707 JP 2003-419273 20031217
PRIORITY APPLN. INFO.: JP 2003-419273 20031217
GI

$$R^2$$
 R^8
 R^7
 R^1
 R^6
 R^8
 R^7
 R^6
 R^8
 R^7
 R^6
 R^8
 R^7
 R^6
 R^8
 R^7

AB The pyrazolylazoquinolines are I (R1-R8 = H, halo, NO2, CN, etc.; R1R2, R3R4, R4R5, R5R6, R6R7, and R7R8 may form ring). The WORM disks, having recording layers containing I-divalent metal chelates, show good heat and light resistance.

MSTR 1

G1 = alkylcarbonylamino (opt. substd.) / CONH2

Patent location: claim 1

Note: additional ring formation also claimed

L5 ANSWER 44 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:86819 MARPAT

TITLE: Colored photoimaging compositions showing good storage

stability for manufacture of color filters

INVENTOR(S):
Kato, Yasuhiro

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 33 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 2005170974 A 20050630 JP 2003-408759 20031208
PRIORITY APPLN. INFO:: JP 2003-408759 20031208

GΙ

$$(R^3)_m$$

$$(R^4)_n$$

$$V$$

$$(R^2)_j$$

$$I$$

$$SO_3Na$$

AB The compns. contain heterocyclic dyes I (R1-R4 = H, substituent; Y = O, N, C; Z = C, N, O, S; when Y = N or C, YZ may form 5- or 6-membered saturated or aromatic ring with C bonded to Y and Nbonded to Z, and ≥1 atoms chosen from C, N, O, and S; when YZ do not form ring, Z = substituent and Y = OH, NHR2, CHR22; m = 0-2; j, n = 0-4). Thus, a pattern from a composition containing

ΙI

II showed good heat and light resistance, and was useful as a color filter for a CCD camera.

MSTR 1

G1 = acylamino / CONH2 / SO2NH2 Patent location: claim 1

L5 ANSWER 45 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 143:16565 MARPAT

TITLE: Azo-substituted quinoline compound and optical

recording material using it

INVENTOR(S): Ueno, Yasunobu; Sato, Tsutomu; Tomura, Tatsuya;

Noguchi, Takashi; Nishimatsu, Masayuki; Maruyama,

Katsuji

PATENT ASSIGNEE(S): Ricoh Co., Ltd., Japan; Chemipro Kasei Ltd.

SOURCE: Jpn. Kokai Tokkyo Koho, 27 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2005146090	A	20050609	JP 2003-384368	20031113
PRIORITY APPLN. INFO.	:		JP 2003-384368	20031113
GI				

Ι

The azo-substituted quinoline compound I (R1-8 = H, halo, nitro, cyano, OH, carboxy, amino, alkyl, aryl, alkyloxy, aryloxy, alkylamino, arylamino, alkylcarbonylamino, arylcarbonylamino, carbamoyl, alkylcarbamoyl, arylcarbamoyl, alkylsulfonylamino, arylsulfonylamino, these may form a ring) and a chelate compound of I and 2-valent metal salt are claimed. Optical recording material comprises a support coated with a recording layer containing the chelate compound The material is suited for high speed recording and large capacity WORM disk.

MSTR 1

G1 = alkylcarbonylamino (opt. substd.) / CONH2

Patent location: claim 1

Note: additional ring formation also claimed

L5 ANSWER 46 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:373698 MARPAT

TITLE: Preparation of 4-aminoquinoline-3-carboxamide

derivatives as PDE4 inhibitors

INVENTOR(S): Edlin, Christopher D.; Eldred, Colin David; Keeling,

Steven Philip; Lunniss, Christopher James; Redfern,

Tracy Jane; Redgrave, Alison Judith; Woodrow, Michael

PATENT ASSIGNEE(S): Glaxo Group Limited, UK SOURCE: PCT Int. Appl., 63 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	PATENT NO. K					DATE			Al		CATI		٥.	DATE			
WO	2005	0302	12	A	1	2005	0407		M				4 4	2004	0923		
	\mathbb{W} :	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	ΒA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
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		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NΙ,
		NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	VN,	YU,	ZA,	ZM,	ZW
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	AZ, BY, I EE, ES, I				KΖ,	MD,	RU,	ΤJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
	EE, ES, 1				FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	ΝL,	PL,	PT,	RO,	SE,
		SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,
		SN,	TD,	ΤG													
EP	1673	086		Α	1	2006	0628		E	P 20	04-7	6565	6	2004	0923		
EP	1673	086		В	1	2008	0123										
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FΙ,	RO,	CY,	TR,	BG,	CZ,	EE,	HU,	PL,	SK,	HR	
	2007																
										T 20	04-7	6565	6	2004	0923		
ES	AT 384530 ES 2298806			T.	3	2008	0516		Ε	S 20	04-7	6565	6	2004	0923		
US	US 20080096884			A	1	2008	0424		U	S 20	07-5	7291	4	2007	0206		
RIORIT	Y APP	LN.	INFO	.:					G1	В 20	03-2	2722		2003	0927		
									M	O 20	04-E	P108	44	2004	0923		
THER S	OURCE	(S):			CAS	REAC'	Г 14:	2 : 37.	3698								

GI

AB The title compds. I [R1 = (un)substituted aryl, heteroaryl, heterocyclyl, etc.; R2 = H, alkyl; R3 = H, alkyl, cycloalkyl, etc.; R4 = H, alkyl; or NR3R4 = (un)substituted heterocyclyl; R5 = H, alkyl; R6 = H, alkyl, alkoxy, etc.] which are inhibitors of phosphodiesterase type IV (PDE4) and are of use in the treatment of inflammatory and/or allergic diseases, were prepared Thus, reacting 4-chloro-6-(1-piperidinylsulfonyl)-3-quinolinecarboxamide (preparation given) with 3-methoxyaniline afforded II. The exemplified compds. I inhibit the catalytic activity at PDE4B (human recombinant) enzyme with pIC50's in the range 7.5-10.8. The pharmaceutical compns. comprising the compound I are disclosed.

Ι

MSTR 1

G1 = Ph (opt. substd. by 1 or more G22)

G19 = morpholino

Patent location: claim 1

Note: or pharmaceutically acceptable salts

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 47 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:355178 MARPAT

TITLE: Preparation of aminocarbonylquinoline derivatives as

phosphodiesterase type IV (PDE4) inhibitors

INVENTOR(S): Edlin, Christopher; Eldred, Colin David; Lunniss,

Christopher James; Redgrave, Alison Judith; Robinson,

John Edward; Woodrow, Michael

PATENT ASSIGNEE(S): Glaxo Group Limited, UK SOURCE: PCT Int. Appl., 69 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	ATENT	NO.		KI	ND	DATE			A.	PPLI	CATI	и ис	0.	DATE			
WC	2005	0307	25	А	1	2005	0407		M	0 20	04-G	B410	6	2004	0927		
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	AZ,	BA,	BB,	ВG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NΙ,
		NO,	NΖ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW: BW, G				KΕ,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,
		ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ΤJ,	TM,	ΑT,	ΒE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,
		SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,
		SN,	TD,	ΤG													
EF	1673	345		Α	1	2006	0628		E.	P 20	04 - 7	6864	9	2004	0927		
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FΙ,	RO,	CY,	TR,	BG,	CZ,	EE,	HU,	PL,	SK,	HR	
JF	JP 2007506717					2007	0322		J.	P 20	06-5	2748.	3	2004	0927		
US	US 20070191426				1	2007	0816		U	S 20	07-5	7291.	3	2007	0206		
PRIORIT	ORITY APPLN. INFO								G:	В 20	03-2	2726		2003	0927		
									M	0 20	04-G	B410	6	2004	0927		
OTHER S	SOURCE	(S) ·			CAS	REAC	т 14	2 • 35	5178								

OTHER SOURCE(S): CASREACT 142:355178

GI

AΒ Title compds. I [R1 = (un)substituted-aryl, -heteroaryl, cycloalkyl, etc.; R2 = H, alkyl; R3 = H, (un)substituted alkyl, cycloalkyl, etc.; R4 = H, alkyl; or R3 and R4 together = (un)substituted N-heterocycle; R5 = H, alkyl; R6 = H, alkoxy, Cl, etc.] and their pharmaceutically acceptable salts, are prepared and disclosed as inhibitors of phosphodiesterase type IV (PDE4). Thus, e.g., II was prepared by amidation of $3-(aminocarbony1)-4-\{[3-$ (methyloxy)phenyl]amino}-6-quinolinecarboxylic acid (preparation given) with morpholine. The inhibition capability of I was evaluated in radioactive Scintillation Proximity Assay (SPA) and revealed that selected compds. of the invention possessed pIC50 values in the range of 6.3-9.5. I as PDE4 inhibitors should prove useful in the treatment of inflammatory and allergic diseases.

MSTR 1

G1 = Ph (opt. substd. by 1 or more G22) = morpholino

Patent location: claim 1 Note: or pharmaceutically acceptable salts

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 48 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:331863 MARPAT

TITLE: Crystal structure of human PIM-1 kinase and use of

structural information for preparation of molecular

scaffolds for kinase ligand development and

pharmaceutical applications

INVENTOR(S): Artis, Dean R.; Bremer, Ryan E.; Gillette, Samuel J.;

Hurt, Clarence R.; Ibrahim, Prabham L.; Zuckerman,

Rebecca L.

PATENT ASSIGNEE(S): Plexxikon, Inc., USA

SOURCE: PCT Int. Appl., 236 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO. KIND DATE
                                        APPLICATION NO. DATE
WO 2005028624 A2 20050331
WO 2005028624 A3 20061102
                                         WO 2004-US30360 20040915
    W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
        CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
        LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
        NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
        TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
    RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
        AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
        EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
        SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
        SN, TD, TG
US 20050164300
                  A1 20050728
                                         US 2004-941635
                                                          20040915
```

PRIORITY APPLN. INFO.:

US 2003-503277P 20030915

AB Mol. scaffolds for compds. active on protein kinases are described, along

Mol. scaffolds for compds. active on protein kinases are described, along with methods for using such scaffolds for kinase ligand development. The use of kinase structural information, exemplified with PIM-1 crystals and structural information can be used for identifying mol. scaffolds and for developing ligands that bind to and modulate particular kinases. More specifically, crystal structure and mol. structural coordinates of human PIM-1 kinase are disclosed. Preparation of compds. modulating PIM-1 and other protein kinases activity (i.e., kinase scaffold library) is reported. These compds. can be used for the treatment of diseases, such as cancer and inflammation.

MSTR 7

```
G11 G11
                  G11
G11
                  G11
      Ġ1
G3
        = 0
G4
        = NH2 (opt. substd.)
G6
        = heteroaryl <containing up to 10 atoms,
           zero or more N, zero or more O,
           zero or more S (no other heteroatoms), mono- or bicyclic>
G10
        = cycloalkyl <containing 3-15 C>
G11
        = 51 / 55 / 56
              _G12-G10
                             02S----G6
G12 = NH
Patent location:
                                    claim 1
Note:
                                    additional substitution also claimed
Note:
                                    substitution is restricted
      ANSWER 49 OF 131 MARPAT COPYRIGHT 2008 ACS on STN
                               142:280200 MARPAT
ACCESSION NUMBER:
TITLE:
                                Preparation of pyrazolylmethylbenzamides as P2X7
                                receptor antagonists
INVENTOR(S):
                                Concepcion, Arnel; Inoue, Tadashi; Mochizuki, Yuki;
                                Muramatsu, Aiko; Gantner, Florian; Nakashima, Kosuke;
                                Urbahns, Klaus; Bacon, Kevin B.
PATENT ASSIGNEE(S):
                                Bayer Healthcare A.-G., Germany
                                PCT Int. Appl., 40 pp.
SOURCE:
                                CODEN: PIXXD2
DOCUMENT TYPE:
                                Patent
LANGUAGE:
                                English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                       APPLICATION NO. DATE
                      KIND DATE
      PATENT NO.
                           ____
                                   _____
                                                       ______
      WO 2005019182 A1 20050303
                                                    WO 2004-EP9172 20040816
           W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
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RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,

AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,

SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,

SN, TD, TG

PRIORITY APPLN. INFO.: EP 2003-18629 20030820

OTHER SOURCE(S): CASREACT 142:280200

GΙ

$$R^3$$
 R^4
 R^2
 HN
 R^1
 I

$$\begin{array}{c|c} & H \\ N \\ \end{array} \begin{array}{c} O \\ Me \\ \end{array}$$

The present invention relates to novel pyrazolylmethylbenzamides I [R1 = $\frac{1}{2}$] AΒ (un) substituted aryl, heteroaryl, alkyl; R2 = alkyl, haloalkyl; R3 = (un) substituted heteroaryl, Ph; R4 = (un) substituted alkyl, alkenyl, etc.], processes for preparing them and pharmaceutical prepns. containing them. Thirty compds. I were prepared E.q., a multi-step synthesis of II, starting from 3-chloromethylbenzoyl chloride and m-anisidine, was given. The pyrazolylmethylbenzamides I exhibit enhanced potency for P2X7 receptor antagonism (no data given) and can be used for the prophylaxis and treatment of diseases associated with P2X7 receptor activity. More specifically, the compds. I are useful for treatment and prophylaxis of diseases as follows: rheumatoid arthritis, osteoarthritis, psoriasis, allergic dermatitis, asthma, chronic obstructive pulmonary disease (COPD), hyperresponsiveness of the airway, septic shock, glomerulonephritis, irritable bowel disease, Crohn's disease, ulcerative: colitis, atherosclerosis, growth and metastases of malignant cells, myoblastic leukemia, diabetes, Alzheimer's disease, meningitis, osteoporosis, burn injury, ischemic heart disease, stroke and varicose veins.

MSTR 1

G1 = quinolinyl (opt. substd. by (1-2) G23) G23 = CONH2 / alkylamino <containing 1-6 C>

Patent location: claim 1

Note: or tautomeric forms, or salts

Stereochemistry: or stereoisomeric forms

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 50 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:261788 MARPAT

TITLE: Preparation of aryl and heteroaryl amino acid

derivatives as antagonists of factor IX and/or factor

XΙ

INVENTOR(S): Mjalli, Adnan M. M.; Andrews, Robert C.; Guo,

Xiao-Chuan; Christen, Daniel Peter; Gohimmukkula, Devi

Reddy; Huang, Guoxiang; Rothlein, Robert; Tyagi, Sameer; Yaramasu, Tripura; Behme, Christopher Transtech Pharma, Inc., USA

PATENT ASSIGNEE(S): Transtech Pharma, Inc., USA SOURCE: PCT Int. Appl., 313 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATE	I TN	. O <i>l</i>		KIND DATE					A.	PPLI	CATI	N NC	0.	DATE			
WO 2		-		 A: A:		2005 2005	-		M	20	04-U	S254	 63	2004	0806		
	W:	AE, CN, GE, LK, NO, TJ, BW, AZ, EE,	AG, CO, GH, LR, NZ, TM, GH, BY, ES, SK,	AL, CR, GM, LS, OM, TN, GM, KG, FI,	AM, CU, HR, LT, PG, TR, KE, KZ,	AT, CZ, HU, LU, PH, TT, LS, MD, GB,	AU, DE, ID, LV, PL, TZ, MW, RU, GR,	DK, IL, MA, PT, UA, MZ, TJ, HU,	DM, IN, MD, RO, UG, NA, TM, IE,	DZ, IS, MG, RU, US, SD, AT, IT,	EC, JP, MK, SC, UZ, SL, BE, LU,	EE, KE, MN, SD, VC, SZ, BG, MC,	EG, KG, MW, SE, VN, TZ, CH,	BY, ES, KP, MX, SG, YU, UG, CY, PL,	FI, KR, MZ, SK, ZA, ZM, CZ, PT,	GB, KZ, NA, SL, ZM, ZW, DE, RO,	GD, LC, NI, SY, ZW AM, DK, SE,
US 2 EP 1	531 0050 0050 6604	2635 796 0049: 0059 439	310 713	A A A A	1 1 1 2	2005 2005 2005 2005 2006	0217 0303 0317 0531		C. U. E.	A 20 S 20 S 20 P 20	04-2 04-2 04-9 04-9 04-7	5317 1388 1321 8031	96 2 6 8	2004 2004 2004 2004 2004 NL,	0806 0806 0806 0806	MC	DТ

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR
CN 1832920 A 20060913 CN 2004-80022750 20040806
JP 2007501844 T 20070201 JP 2006-523245 20040806
PRIORITY APPLN. INFO.: US 2003-493878P 20030808
US 2003-493879P 20030808
US 2003-493903P 20030808
WO 2004-US25463 20040806

OTHER SOURCE(S): CASREACT 142:261788

The invention relates to aryl and heteroaryl compds. Ar2-K [Ar2 is (un) substituted aryl, heteroaryl, fused cycloalkylaryl, fused cycloalkylheteroaryl, fused heterocyclylaryl or fused heterocyclylheteroaryl; K is a carbamoyl group of defined structure or Ar1-V-CH[(CH2)0-2-G]-X-, where G is H, CO2R1, CH2OR1, COR1, CR1:NOR2, CONR1R2, CONHNH2 or an acid or ester isostere and R1, R2 independently are H, alkyl, alkoxy, aryl, alkylaminoacyl, etc. or may combine to form a ring; V is (CH2)1-2-S-(CH2)0-2, (CH2)1-2-S, S-(CH2)0-2 (or corresponding sulfonyl derivs.), (CH2)1-2-0-(CH2)0-2, (CH2)1-2-NR7-(CH2)0-2, (CH2)1-2-0or a direct bond, where R7 is H, alkyl, aryl, etc. (the CH2 or CH2CH2 groups may be substituted); X is NR8, CONR8, NR8CO, NR8CONR9, O2CNR8, SO2NR8 or NR8SO2NR9, where R8, R9 are independently H, alkyl, aryl, etc.; Ar1 is a group as defined for Ar2] and their pharmaceutical compns. Compds. Ar2-K may be antagonists or partial antagonist of factor IX and/or factor XI and thus may be useful for inhibiting the intrinsic pathway of blood coagulation. Applications include the management, treatment and/or control of diseases caused in part by the intrinsic clotting pathway. Thus, (25)-[5-bromo-2-(4-trifluoromethylbenzyloxy)benzoylamino]-3-(2'phenoxybiphenyl-4-yl)propionic acid, prepared by amidation and O-benzylation reactions, inhibited factor IX or factor XI in the in vitro clotting assay with IC50 < 30 micromolar.

MSTR 1

$$G1 = 564$$

G36 = 577 / 580 / 585

G37 = N / 567

C——G36 567

G41 = NH G45 = NH2

Patent location: claim 1

Note: additional derivatization also claimed

L5 ANSWER 51 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:170068 MARPAT

TITLE: Small molecule toll-like receptor (TLR) antagonists

INVENTOR(S): Lipford, Grayson B.; Forsbach, Alexandra; Zepp,

Charles M.

PATENT ASSIGNEE(S): Coley Pharmaceutical G.m.b.H., Germany; Coley

Pharmaceutical Group, Inc.

SOURCE: PCT Int. Appl., 193 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

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PATENT NO. KIND DATE
                                       APPLICATION NO. DATE
    _____
                                         _____
    WO 2005007672 A2 20050127
WO 2005007672 A3 20050915
                                         WO 2004-US19714 20040618
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
            CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
            GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
            LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
            NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
            TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
        RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
            AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
            EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
            SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
            SN, TD, TG
                    A1
    AU 2004257149
                         20050127
                                         AU 2004-257149
                                                         20040618
    CA 2528774
                    A1 20050127
                                       CA 2004-2528774 20040618
                    A1
    US 20050119273
                         20050602
                                         US 2004-872196
                                                         20040618
    US 7410975
                    В2
                          20080812
    EP 1635846
                    A2 20060322
                                         EP 2004-776820 20040618
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR
                                         CN 2004-80017064 20040618
    CN 1809357
                    A 20060726
    BR 2004011514
                          20060801
                                         BR 2004-11514
                                                       20040618
                                                        20040618
                    Τ
    JP 2007524615
                          20070830
                                         JP 2006-517471
                   A 20060224
A1 20071004
A 20070706
                                         MX 2005-PA13922 20051216
    MX 2005PA13922
    US 20070232622
                                         US 2006-543314
                                                          20061004
    IN 2006KN00153
                                         IN 2006-KN153
                                                          20061119
                                         US 2003-480588P 20030620
PRIORITY APPLN. INFO.:
                                         US 2004-556007P 20040323
                                         US 2004-872196
                                                         20040618
                                         WO 2004-US19714 20040618
```

AB The invention provides methods and compns. useful for modulating signaling through Toll-like receptors (TLR). The methods involve contacting a TLR-expressing cell with a small mol. having a core structure including at least two rings. Certain of the compds. are 4-primary amino quinolines. Many of the compds. and methods are useful specifically for inhibiting immune stimulation involving at least one of TLR9, TLR8, TLR7, and TLR3. The methods may have use in the treatment of autoimmunity, inflammation, allergy, asthma, graft rejection, graft vs. host disease, infection, sepsis, cancer, and immunodeficiency.

MSTR 8

G6 = CONH2 G7 = 35-7 36-34

G8—G9

G8 = NH

G9 = alkylene < containing 1-10 C>

G17 = SO2NH2

Patent location: claim 89

Note: additional ring formation also claimed

Note: or pharmaceutically acceptable hydrates or salts

L5 ANSWER 52 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:134612 MARPAT

TITLE: Preparation of 4-arylaminoquinazolines and analogs as

activators of caspases and inducers of apoptosis INVENTOR(S): Cai, Sui Xiong; Sirisoma, Nilantha Sudath; Pervin,

Azra; Drewe, John A.; Kasibhatla, Shailaja; Jaing, Songchun; Zhang, Hong; Pleiman, Chris; Baichwal,

Vijay; Manfredi, John; Bhoite, Leena

PATENT ASSIGNEE(S): Myriad Genetics, Inc., USA; Cytovia, Inc.

SOURCE: PCT Int. Appl., 289 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2005003100 WO 2005003100	A2 A3	20050113 20050512	WO 2004-US21631	20040706

GI

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
             EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
             SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
             SN, TD,
                     ΤG
                                            AU 2004-253967
     AU 2004253967
                       Α1
                            20050113
                                                             20040706
     CA 2531327
                            20050113
                                            CA 2004-2531327
                                                             20040706
                       Α1
     EP 1660092
                       Α2
                            20060531
                                            EP 2004-785803
                                                             20040706
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR
     CN 1984660
                            20070620
                                            CN 2004-80024205 20040706
                       Α
     JP 2007524637
                       Τ
                            20070830
                                            JP 2006-517854
                                                             20040706
     IN 2006KN00019
                       Α
                            20070316
                                            IN 2006-KN19
                                                             20060102
                                            US 2003-484325P
PRIORITY APPLN. INFO.:
                                                             20030703
                                            US 2003-493006P
                                                             20030807
                                            US 2004-557556P
                                                             20040329
                                            US 2004-571288P
                                                             20040514
                                            WO 2004-US21631
                                                             20040706
```

AΒ 4-Arylaminoquinazolines and analogs I [wherein A = 6-membered (hetero)aryl or carbocycle; L = [C(RL1)(RL2)]n or -N(RL1)C(0)-; RL1, RL2 = H or alkyl;n = 0-2; R1 = Me or ethyl; Ar = (un) substituted (hetero) aryl; R2-R6, R12-R17 = H, halo, N3, OH, thiol, nitro, CN, NH2, alk(en/yn)yl or alkoxy; B, D, Q, T, U, V = C or N, wherein at least one of B and D is N; etc. or pharmaceutically acceptable salts or solvates thereof] were prepared as activators of caspases and inducers of apoptosis. For example, 2,4-quinazolinedione was refluxed with neat phosphorylchloride to give 2,4-dichloroquinazoline in 96% yield, which was coupled with 4-methoxy-N-methylaniline to afford II in 87% yield. II exhibited caspase activation (EC50 2 nM for human breast cancer cell line T-47D, 24 h), inhibition of cell proliferation (GI50 8 nM for T-47D), inhibition of tubulin polymerization (IC50 <500 nM) and cytotoxicity in multidrug resistant cells (IC50 2.9 nM for MCF-7 cell line). Other biol. activities of the invented compds. have also been tested. Therefore, I and pharmaceutical compns. thereof (examples given) are effective activators of caspases and

inducers of apoptosis, and useful in the treatment of such as cancer, autoimmune and inflammation. Disclosed are 4-arylaminoquinazolines and analogs thereof effective as activators of caspases and inducers of apoptosis.

MSTR 1

```
G1
       = CH=CHCH=CH (opt. substd. by G2)
G2
2<sup>C</sup> (0)-G8
     = NH2 / piperidino
       = 1 \text{ or more N} / 31
     -G10
G10
     = 46
₄Ç(O)-G8
G11
     = Me
G12
     = G13
G13
       = (0-3) CH2
       = Ph (opt. substd. by G17)
```

L5 ANSWER 53 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 142:23205 MARPAT

TITLE: Preparation of quinoline derivatives as

claim 1

phosphodiesterase inhibitors

INVENTOR(S): Baldwin, Ian Robert; Barker, Michael David; Dean,

Anthony William; Eldred, Colin David; Evans, Brian; Gough, Sharon Lisa; Guntrip, Stephen Barry; Hamblin, Julie Nicole; Holman, Stuart; Jones, Paul; Lindvall, Mika Kristian; Lunniss, Christopher James; Redfern, Tracy Jane; Redgrave, Alison Judith; Robinson, John

Edward; Woodrow, Michael Glaxo Group Limited, UK

PATENT ASSIGNEE(S): Glaxo Group Limited, UK SOURCE: PCT Int. Appl., 243 pp.

Patent location:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	TENT	NO.		KI	ND	DATE			A	PPLI	CATI	N NC	Ο.	DATE			
WO	2004 W: RW:	AE, CN, GE, LK, NO, TJ, BW, AZ, EE, SI,	AG, CO, GH, LR, NZ, TM, GH, BY, ES,	CR, GM, LS, OM, TN, GM, KG, FI, TR,	AM, CU, HR, LT, PG, TR, KE, KZ,	CZ, HU, LU, PH, TT, LS, MD, GB,	AU, DE, ID, LV, PL, TZ, MW, RU, GR,	AZ, DK, IL, MA, PT, UA, MZ, TJ,	BA, DM, IN, MD, RO, UG, NA, TM, IE,	DZ, IS, MG, RU, US, SD, AT, IT,	BG, EC, JP, MK, SC, UZ, SL, BE, LU,	BR, EE, KE, MN, SD, VC, SZ, BG, MC,	BW, EG, KG, MW, SE, VN, TZ, CH,	2004 BY, ES, KP, MX, SG, YU, UG, CY, PL, GW,	BZ, FI, KR, MZ, SK, ZA, ZM, CZ, PT,	GB, KZ, NA, SL, ZM, ZW, DE, RO,	GD, LC, NI, SY, ZW AM, DK, SE,
CA EP	2004 2526 1633 1633	2407 228 748		A: A: A: B:	1 1	2004 2004 2006 2008	1202 0315		C.	U 20 A 20 P 20	04-2	5262	28	2004 2004 2004	0519		
CN JP AT ES	R: 2004 1823 2007 3881 2301 1944	IE, 0104 063 5012 48 993	SI, 77		LV,		RO, 0530 0823 0125 0315 0701	CY,	TR, B: C: J: A E	BG, R 20 N 20 P 20 T 20	CZ, 04-1 04-8 06-5 04-7	EE, 0477 0020 2988 3379 3379	HU, 651 9 9	NL, PL, 2004 2004 2004 2004 2004 2004	SK, 0519 0519 0519 0519 0519		PT,
NO US MX IN US	R: 2005 2007 2005 2006 2006	AT, IT, 0054 0142 PA12 KN02 0178 0049	LI, 21 373 466 416 416 570	BG, LU, A A A A	CH, MC,	CY,	CZ, PL, 1220 0621 0130 1013 0810	DE, PT,	DK, RO, No U M M II U U GG GG GG EE	EE,	ES, SI, 05-5 05-5 05-P. 06-3 06-3 03-1 03-2 04-7	FI, SK, 421 5707 A124 N241 4967 4970 1688 6187 3379 P549	FR, TR, 9 66 6 7 1	GB, HR, 2005 2005 2005 2006 2006 2006 2003 2003 2004 2004 2004	GR, LT, 1116 1117 1118 1129 0208 0208 0521 1110 0519 0519		IE,
GI																	

Page 103

AB Title compds. represented by the formula I [wherein R1 = (un)substituted (cyclo)alkyl, (hetero)aryl, cycloalkylalkyl, etc.; R2 = H or alkyl; R3 = H, (un)substituted SOnalkyl, 2-oxopyrrolidin-1-yl, cycloalkyl, etc.; R4 = H or SOnalkyl; R5 = H, halo, alkyl, alkoxy; n = 0-2; and pharmaceutically acceptable salts thereof] were prepared as phosphodiesterase inhibitors. For example, reaction of 4-chloro-6-(methylsulfonyl)-3-quinolinecarboxamide with 3-fluoroaniline gave II. Selected prepared compds. were tested for inhibition of PDE4B (human recombinant) enzyme and PDE5 with pIC50 values in the range of 6.0-11.7 and 4.5-7.0, resp. Thus, I and their pharmaceutical compns. are useful as phosphodiesterase inhibitors, especially PDE4 inhibitors, for the prophylaxis or treatment of a clin. condition, such as inflammatory and/or allergic diseases (no data).

MSTR 1

G1 = benzothiazolyl

G8 = NHG22 = 84

G23-G24

G23 = S02

G24 = piperidino (substd. by 1 or more 335)

335 G(O)—G50

G43 = 146

G23—G24 146

G55 = 11

G1 11

Patent location: claim 1

Note: also incorporates claim 25 structures II, III, and

XXIX

Note: substitution is restricted

Note: additional oxo formation also claimed Note: or pharmaceutically acceptable salts

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 54 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 141:314346 MARPAT

TITLE: Preparation of quinoline, tetrahydroquinazoline, and

pyrimidine derivatives as MCH antagonist for treatment

of CNS disorders

INVENTOR(S): Sekiguchi, Yoshinori; Kanuma, Kosuke; Omodera,

Katsunori; Busujima, Tsuyoshi; Tran, Thuy-Anh; Han, Sangdon; Casper, Martin; Kramer, Bryan A.; Semple,

Graeme; Zou, Ning

PATENT ASSIGNEE(S): Taisho Pharmaceutical Co. Ltd., Japan; Arena

Pharmaceuticals, Inc.

SOURCE: Eur. Pat. Appl., 586 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
EP 1464335	A2 20041006	EP 2004-7651	20040330
EP 1464335	A3 20070509		
R: AT, BE,	CH, DE, DK, ES, FI	k, GB, GR, IT, LI, LU	, NL, SE, MC, PT,
IE, SI,	LT, LV, FI, RO, MI	I, CY, AL, TR, BG, CZ	, EE, HU, PL, SK
US 20050197350	A1 20050908	US 2004-812075	20040330
AU 2004226049	A1 20041014	AU 2004-226049	20040331
CA 2518913	A1 20041014	CA 2004-2518913	20040331
WO 2004087669	A1 20041014	WO 2004-JP4624	20040331
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NO, NZ,	OM, PG, PH, PL, PT	C, RO, RU, SC, SD, SE	, SG, SK, SL, SY,
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RW: BW, GH,	GM, KE, LS, MW, M	i, SD, SL, SZ, TZ, UG	, ZM, ZW, AM, AZ,
BY, KG,	KZ, MD, RU, TJ, Ti	I, AT, BE, BG, CH, CY	, CZ, DE, DK, EE,

ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG JP 2004300156 20041028 JP 2004-107965 20040331 Α BR 2004008910 Α 20060321 BR 2004-8910 20040331 CN 2004-80014547 20040331 CN 1798736 Α 20060705 IN 2005KN01805 Α 20061201 IN 2005-KN1805 20050912 MX 2005PA10475 Α 20060525 MX 2005-PA10475 20050929 NO 2005004999 Α 20051107 NO 2005-4999 20051027 PRIORITY APPLN. INFO.: US 2003-458530P 20030331 US 2003-495911P 20030819 US 2003-510186P 20031009 US 2003-530360P 20031216 WO 2004-JP4624 20040331 GΙ

$$(T)_{p} \xrightarrow{\mathbb{R}^{2}} (T)_{p} \xrightarrow{\mathbb{R}^{2}} \mathbb{N}$$

$$(T)_{p} = \begin{bmatrix} R^{2} \\ N \\ N \end{bmatrix}$$

AB Title compds. I, II, and III [wherein R1 = (un)substituted (cyclo)alkyl, (cyclo)alkenyl, alkynyl, aryl; R2 = H, halo, OH, carboxy, carbamoyl, amino, (un)substituted alkyl, alkoxy; T = independently H, halo, OH, carboxy, carbamoyl, amino, cyano, NO2, alkenyl, alkynyl, cycloalkyl, (un)substituted alkyl, alkoxy; p = 0-5; L = aminocycloalkylideneamino, etc.; Y = bond, CH2, CO2, OCO, SO2, CO, CS, CONH, CSNH, etc.; with provisos; and pharmaceutically acceptable salts, hydrates, or solvates thereof] were prepared as antagonists of melanin concentrating hormone (MCH),

an

endogenous ligand of G-protein coupled receptors (GPCRs). Examples include solution and solid phase general synthetic methods and phys. data for nearly 3400 invention compds. In addition, all exemplified compds. were assayed using high throughput functional screening to detect intracellular Ca2+ concns. for accessing GPCR activation. For instance, reaction of 2,4-dichloro-6-methylpyrimidine with dimethylamine gave 2-chloro-4-(dimethylamino)-6-methylpyrimidine (40%), which was coupled with cis-(4-aminocyclohexyl)carbamic acid tert-Bu ester (60%). Deprotection (72%), amidation, and workup provided the benzamide ${
m IV}ullet{
m TFA}$. The latter demonstrated MCH antagonist activity with an IC50 value of 7.6 nM. Thus, pharmaceutical compns. comprising I are useful for the prophylaxis or treatment of improving memory function, sleeping and arousal, anxiety, depression, mood disorders, seizure, obesity, diabetes, appetite and eating disorders, cardiovascular disease, hypertension, dyslipidemia, myocardial infarction, binge eating disorders including bulimia, anorexia, mental disorders including manic depression, schizophrenia, delirium, dementia, stress, cognitive disorders, attention deficit disorder, substance abuse disorders, and dyskinesias including Parkinson's disease, epilepsy, and addiction (no data). This is part I of three in a series covering the patent.

MSTR 1A

G1 = 12-5 14-2

G2 = NHNH2G6 = CONH2

Patent location: claim 1

Note: substitution is restricted

Note: additional substitution also claimed

L5 ANSWER 55 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 141:190691 MARPAT

TITLE: Preparation of heteroaryl amines, in particular

quinolin-4-yl amines, as antagonists for α -2,

especially α -2C, adrenoceptors

INVENTOR(S): Hoeglund, Iisa; Koivisto, Ari-Pekka; Tauber, Andrei;

Kallatsa, Oili; Sallinen, Jukka; Silver, Satu;

Hoffren, Anna-Marja; Iles, Matthew; Wurster, Siegfried

10/572,914

PATENT ASSIGNEE(S): Oy Juvantia Pharma Ltd., Finland

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2004067513 A1 20040812 WO 2004-F138 20040127

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI PRIORITY APPLN. INFO::

FI 2003-120 20030127

US 2003-442570P 20030127

GΙ

Title compds. I [wherein Q = (un)substituted 1,4-phenylene, II or III; R5 = independently OH, halo, alkyl, alkenyl, alkoxy, NO2, etc.; r = 0-2; L =CH, CR5, N; Y = -CHa(R4)d[CHb(R4)c]v or a single bond; R1 = H, cyclo/alkyl; A = benzene ring or (C3-C7)cycloalkyl; each R2 = independently OH, halo, alkenyl, alkynyl, alkyl, alkoxy, NO2, monoalkyl/dialkyl/amino, -S-alkyl, -CO-NH2, CHO, etc.; R3 = H, alkyl, alkenyl, alkylcarbonyl, aminocarbonyl, (un)substituted Ph, naphthyl, benzyl, etc.; R4 = independently OH, halo, amino, oxo, CHO, alkyl, alkenyl, alkynyl, alkylcarbonyl, aminocarbonyl, (un)substituted cycloalkyl, Ph, naphthyl, benzyl, etc.; or R3 and R4 or R4 and R4 with any of the ring atom(s) to which they are attached = condensed (un)substituted 5-7 carbocyclic to heterocyclic ring; Ra, Rb = independently H, OH, halo, alkyl, alkenyl, alkynyl, alkoxy, NO2, monoalkyl/dialkyl/amino, -S-alkyl, CN, (un)substituted cycloalkyl, Ph or 5-6 membered heterocyclyl, etc.; or Rb as defined above and RaCCNR1 = condensed (un)substituted 5-7 membered heterocycle; or RaCCRb = condensed (un)substituted 5-7 membered non-aromatic carbo- or heterocyclic ring; a, b, c, d = independently 0-2; n = 0-3; q =0-4; v = 0-1; with provisos; their pharmaceutically acceptable salts and esters] were prepared as alpha-2, in particular selective α -2C, adrenoreceptor antagonists. Amination of 2-methylpiperidine with 1-chloro-4-nitrobenzene, methylation with MeI, and reduction of the nitro intermediate gave 3-Methyl-1-(4-nitrophenyl)piperazine (IV). Cyclocondensation of 2,3-dimethylaniline with Et 2-methylacetoacetate, chlorination with SO2Cl2, and alkylation of amine IV with the resulting chloride gave the dialkylated amine V. I are useful for treating CNS disorders, especially depression.

MSTR 1

^{*} STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

= p-C6H4 (opt. substd. by (1-3) G2)

= CH=CHCH=CH (opt. substd. by 1 or more G17) G16

G17 = alkylaminocarbonyl <containing 1-6 C>

G18 = CONH2

Patent location: claim 1

Note: or pharmaceutically acceptable salts or esters

Note: substitution is restricted

ANSWER 56 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

141:123658 MARPAT ACCESSION NUMBER:

TITLE: Preparation of antidepressant arylpiperazine derivatives of heterocycle-fused benzodioxans

INVENTOR(S): Evrard, Deborah Ann; Zhou, Dahui; Stack, Gary Paul;

Venkatesan, Aranapakam Madumbai; Failli, Amedeo A.;

Croce, Susan Christman

Wyeth, USA PATENT ASSIGNEE(S):

SOURCE: U.S. Pat. Appl. Publ., 30 pp., Cont.-in-part of U.S.

Provisional Ser. No. 410,082.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT NO.		APPLICATION NO. DATE
US 20040142926 US 7153849		2 US 2003-659537 20030910
CA 2497783	A1 2004032	5 CA 2003-2497783 20030911
WO 2004024731	A1 2004032	5 WO 2003-US28453 20030911
W: AE, AG,	AL, AM, AT, A	, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
CO, CR,	CU, CZ, DE, DI	, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE,
GH, GM,	HR, HU, ID, II	, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
LR, LS,	LT, LU, LV, M	, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM,
PG, PH,	PL, PT, RO, RI	, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
TR, TT,	TZ, UA, UG, US	, UZ, VC, VN, YU, ZA, ZM, ZW
RW: GH, GM,	KE, LS, MW, M	, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
KG, KZ,	MD, RU, TJ, Ti	, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
FI, FR,	GB, GR, HU, II	, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
BF, BJ,	CF, CG, CI, CI	, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
AU 2003272316	A1 2004043	0 AU 2003-272316 20030911
EP 1537121	A1 2005060	8 EP 2003-754492 20030911
R: AT, BE,	CH, DE, DK, ES	, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI,	LT, LV, FI, R	, MK, CY, AL, TR, BG, CZ, EE, HU, SK

BR	2003014277	A	20050726	BR	2003-14277	20030911
CN	1681822	A	20051012	CN	2003-821677	20030911
JP	2006507250	Τ	20060302	JP	2004-536475	20030911
CN	101239953	A	20080813	CN	2007-10142627	20030911
MX	2005PA02743	A	20050603	MX	2005-PA2743	20050311
US	20060276481	A1	20061207	US	2006-505663	20060816
PRIORIT	Y APPLN. INFO.:			US	2002-410082P	20020912
				US	2003-659537	20030910
				CN	2003-821677	20030911
				WO	2003-US28453	20030911

Ι

GΙ

The title compds. [I; R1 = H, halo, CN, carboxamido, etc.; XY = AB N:CR2CR3:N, N:CR2CR4:CH, N:CR2N:CH, N:CR2O, NHCR5:CH; R2, R3 = H, halo, NH2, mono- or dialkylamino, alkyl; R4 = H, alkyl; R5 = H, halo, CF3, pentafluoroethyl, alkyl; Ar = (un)substituted Ph, naphthyl, indolyl, indazolyl, thienyl, etc.; n = 1-2], useful for the treatment of depression (including but not limited to major depressive disorder, childhood depression and dysthymia), anxiety, panic disorder, post-traumatic stress disorder, premenstrual dysphoric disorder (also known as premenstrual syndrome), attention deficit disorder (with and without hyperactivity), obsessive compulsive disorder, social anxiety disorder, generalized anxiety disorder, obesity, eating disorders such as anorexia nervosa and bulimia nervosa, vasomotor flushing, cocaine and alc. addiction, sexual dysfunction and related illnesses, were prepared Thus, reacting [(2S)-8-methyl-2,3-dihydro[1,4]dioxino[2,3-f]quinolin-2-yl]methyl4-bromobenzenesulfonate with 3-chlorophenylpiperazine. HCl in the presence of EtN(iso-Pr)2 in DMSO afforded 68% (2S)-2-{[4-(3-chlorophenyl)piperazin-1-y1]methyl}-8-methyl-2,3-dihydro[1,4]dioxino[2,3-f]quinoline. The exemplified compds. I were tested for 5-HT transporter affinity, 5-HT1A receptor affinity, and antagonistic activity at 5-HT1A receptors and biol. data were given. The pharmaceutical composition comprising the compound I is claimed.

MSTR 1

= quinolinyl (opt. substd. by (1-3) G14) = CONH2 / alkylamino <containing 1-6 C>

Patent location: claim 1

Note: or pharmaceutically acceptable salts

REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 57 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

140:339123 MARPAT ACCESSION NUMBER:

TITLE: Preparation of podophyllotoxin derivatives as

anticancer compounds

INVENTOR(S): Shi, Qian; Wang, Hui-kang; Oyama, Masayoshi; Vance,

John Robert; Chen, Ming S.

Plantaceutica Inc., USA PATENT ASSIGNEE(S): PCT Int. Appl., 52 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

```
PATENT NO. KIND DATE
                                         APPLICATION NO. DATE
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    WO 2004033423 A2 20040422
WO 2004033423 A3 20040729
                                         WO 2003-US32547 20031014
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE,
            GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
            LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,
            OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
            TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
            KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
            FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                       CA 2003-2501901 20031014
    CA 2501901
                    A1 20040422
    AU 2003300385
                    A1 20040504
                                         AU 2003-300385 20031014
                    A1
                         20040715
                                         US 2003-685870
    US 20040138288
                                                         20031014
    US 6903133
                     В2
                          20050607
    EP 1610790
                    A2 20060104
                                         EP 2003-808232
                                                        20031014
           AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
    JP 2006503079 T 20060126
                                         JP 2004-543785 20031014
PRIORITY APPLN. INFO.:
                                         US 2002-417785P 20021011
                                         WO 2003-US32547 20031014
```

GΙ

AB Podophyllotoxin derivs., such as I [R1, R2, R3, R7 = H, alkyl; R4, R6 = alkyl; R5 = H, P(0) (ORa)2; Ra = H, alkyl; T = H; XT = :N; X = bond, O, S, NRb; Rb = H, alkyl; Y = 5-membered heteroaryl or heterocyclyl, optionally substituted with one or more halogen, alkyl, cyclyl, aryl, heteroaryl, heterocyclyl, etc.], were prepared for their therapeutic use as anticancer agents. Thus, podophyllotoxin derivative II was prepared via a multistep synthetic sequence starting from 4'-demethyl-4 β -bromo-4-desoxypodophyllotoxin (prepared from podophyllotoxin), 2-aminothiazole-4-acetic acid and (trimethylsilyl)diazomethane. II showed unexpectedly high levels of cellular protein-linked DNA breaks (PLDB) induction in KB cells when tested at $5\mu g/mL$. This invention also features a method for treating cancer.

MSTR 1

 $G2 = 48-8 \ 49-117 \ 48-1$

G8 = NH

G9 = quinolinyl (opt. substd. by G27)

G27 = 218

C(O)-G29

G29 = NH2

Patent location: claim 1

L5 ANSWER 58 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 140:287410 MARPAT

TITLE: Preparation of antidepressant arylpiperazine

derivatives of heterocycle-fused benzodioxans

INVENTOR(S): Evrard, Deborah A.; Zhou, Dahui; Stack, Gary Paul;

Venkatesan, Arenapakam Madumbai; Failli, Amedeo A.;

Croce, Susan Christman

PATENT ASSIGNEE(S): Wyeth, John, and Brother Ltd., USA

SOURCE: PCT Int. Appl., 84 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

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APPLICATION NO. DATE
    PATENT NO. KIND DATE
                                    WO 2003-US28453 20030911
    WO 2004024731 A1 20040325
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE,
            GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
            LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM,
            PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
            TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
            KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
            FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                        US 2003-659537 20030910
    US 20040142926
                    A1 20040722
    US 7153849
                     B2 20061226
    CA 2497783
                     A1
                         20040325
                                         CA 2003-2497783 20030911
    AU 2003272316 A1 20040430
EP 1537121 A1 20050608
                                         AU 2003-272316
                                                         20030911
                                    EP 2003-754492 20030911
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
    BR 2003014277 A 20050726
JP 2006507250 T 20060302
                                         BR 2003-14277 20030911
                                          JP 2004-536475
                                                         20030911
                    А
                                          MX 2005-PA2743 20050311
    MX 2005PA02743
                        20050603
                                          US 2002-410082P 20020912
PRIORITY APPLN. INFO.:
                                          US 2003-659537
                                                          20030910
                                          WO 2003-US28453 20030911
```

GΙ

AΒ The title compds. [R1 = H, halo, CN, carboxamido, etc.; XY = N:CR2CR3:N, N:CR2CR4:CH, N:CR2N:CH, N:CR2O, NHCR5:CH; R2, R3 = H, halo, NH2, mono-or dialkylamino, alkyl; R4 = H, alkyl; R5 = H, halo, CF3, pentafluoroethyl, alkyl; Ar = (un)substituted Ph, naphthyl, indoleyl, indazolyl, thienyl, etc.; n = 1-2], useful for the treatment of depression (including but not limited to major depressive disorder, childhood depression and dysthymia), anxiety, panic disorder, post-traumatic stress disorder, premenstrual dysphoric disorder (also known as premenstrual syndrome), attention deficit disorder (with and without hyperactivity), obsessive compulsive disorder, social anxiety disorder, generalized anxiety disorder, obesity, eating disorders such as anorexia nervosa and bulimia nervosa, vasomotor flushing, cocaine and alc. addiction, sexual dysfunction and related illnesses, were prepared Thus, reacting [(2S)-8-methyl-2,3dihydro[1,4]dioxino[2,3-f]quinolin-2-yl]methyl 4-bromobenzenesulfonate with 3-chlorophenylpiperazine. HCl in the presence of EtN(iso-Pr)2 in DMSO 2,3-dihydro[1,4]dioxino[2,3-f]quinoline. The exemplified compds. I were tested for 5-HT transporter affinity, 5-HT1A receptor affinity, and antagonistic activity at 5-HT1A receptors and biol. data were given. pharmaceutical composition comprising the compound I is claimed.

Ι

MSTR 1

G3 = quinolinyl (opt. substd. by (1-3) G14) G14 = CONH2 / alkylamino <containing 1-6 C>

Patent location: claim 1

Note: or pharmaceutically acceptable salts Note: or pharmaceutically acceptable salts

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 59 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 140:128289 MARPAT

TITLE: Preparation of 8-hydroxyquinolines for treatment of

neurological conditions.

INVENTOR(S): Barnham, Kevin Jeffrey; Gautier, Elisabeth Colette

Louise; Kok, Gaik Beng; Krippner, Guy PATENT ASSIGNEE(S): Prana Biotechnology Limited, Australia

SOURCE: PCT Int. Appl., 149 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

						KIND DATE								DATE				
;	 WО	2004	 0074												2003	0716		
		W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MΖ,	NI,	NO,	NZ,	OM,
			PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ΤJ,	TM,	TN,
			TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW			
		RW:	GH,	GM,	KΕ,	LS,	MW,	MΖ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
			KG,	KΖ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
			FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,
			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	ΤG
	CA	2493	536		Α	1	2004	0122		C.	A 20	03-2	4935	36	2003	0716		
	ΑU	2003	2438	36	Α	1	2004	0202		A	U 20	03-2	4383	6	2003	0716		
		1539								E	P 20	03-7	6351	6	2003	0716		
		R:													NL,		,	PT,
															EE,		SK	
	BR	2003	0129	34	А		2005	0621		B	R 20	03-1	2934		2003	0716		
		1681																
	JΡ	2006	5046	46	Т		2006	0209		J:	P 20	04 - 5	2019	5	2003	0716		
		5376					2007	1026		N	Z 20	03-5	3767	7	2003	0716		
		2005													2005			
		2005			А										2005			
		2006																
		2006																
		2008				1	2008	0703							2007			
PRIOR	IT	APP:	LN.	INFO	.:										2002			
															2003			
															2005			
										U	S 20	05-5	2190	2	2005	0810		
GI																		

$$R^4$$
 R^3
 R
 R^5
 R^2
 R^2
 R^3

AB A method for the treatment of a neurol. condition comprises administration of title compds. [I; R1 = H, (substituted) alkyl, alkenyl, acyl, aryl, heterocyclyl, antioxidant or targeting moiety; R2 = H; (substituted)

alkyl, alkenyl, aryl, heterocyclyl, alkoxy, antioxidant, targeting moiety, COR6, CSR6, etc.; R6 = H, (substituted) alkyl, alkenyl, aryl, heterocyclyl, etc.; R, R', R3, R4, R5 = H, OH, halo, SO3H, cyano, CF3, (substituted) alkyl, alkenyl, alkoxy, acyl, amino, thio, sulfonyl, sulfinyl, sulfonylamino, aryl, heterocyclyl, antioxidant or targeting moiety; with provisos]. Thus, 5,7-dichloro-8-hydroxyquinoline-2-carboxylic acid (preparation given), dicyclohexylcarbodiimide, 1-hydroxybenzotriazole hydrate, histamine dihydrochloride, and Et3N were stirred in DMF/CH2Cl2 to give 34% 5,7-dichloro-8-hydroxyquinoline-2-carboxylic acid [2-(1H-imidazol-4-yl)ethyl]amide (PBT 1038). This inhibited metal-mediated lipoprotein oxidation with IC50 = 0.26 $\mu \rm M$.

MSTR 1

G14 = CONH2 (opt. substd.) / 62

Patent location: claim 1

Note: or salts, hydrates, solvates, derivatives,

prodrugs, tautomers

Note: substitution is restricted

Stereochemistry: or isomers

REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 60 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 140:117442 MARPAT

TITLE: Pharmaceutical compositions comprising hepatitis C

viral protease inhibitors

INVENTOR(S): Chen, Shirlynn; Mei, Xiaohui

PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 66 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004009121	A1	20040129	WO 2003-US22434	20030717

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
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             PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
             TR, TT, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
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             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     US 20040033959
                     A1 20040219
                                          US 2003-620408
                                                          20030716
                                          AU 2003-259155
     AU 2003259155
                      Α1
                            20040209
                                                           20030717
PRIORITY APPLN. INFO.:
                                          US 2002-397280P 20020719
                                          WO 2003-US22434 20030717
```

AB Disclosed are pharmaceutical compns. of hepatitis C viral protease inhibitors having improved bioavailability, and methods of using these compns. for inhibiting the replication of the hepatitis C virus (HCV) and for the treatment of an HCV infection. These compns. include co-solvent systems, lipid based systems, solid dispersions and granulations, and all comprise the hepatitis C viral protease inhibitor, at least one pharmaceutically acceptable amine and optionally one or more addnl. ingredients.

MSTR 1

G20 = 120

G21 = 25 / 27 / 31

G23 = NH

G24 = aryl < containing 6-10 C >

G26 = NH2 / 34

G23-G24

Patent location: claim 1
Note: or tautomers

Note: additional substitution also claimed

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 61 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:381510 MARPAT

TITLE: Preparation of piperazine derivatives as antiviral

agents

INVENTOR(S): Wang, Tao; Wallace, Owen B.; Meanwell, Nicholas A.;

Kadow, John F.; Zhang, Zhongxing; Yang, Zhong

PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA

SOURCE: PCT Int. Appl., 121 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO. KIND DATE
                                       APPLICATION NO. DATE
    WO 2003092695 A1 20031113 WO 2003-US8893 20030321
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
            PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT,
            TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
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            FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    US 20040009985
                   A1 20040115
                                       US 2003-393030
                                                       20030320
    US 7037913
                    В2
                        20060502
    AU 2003220480
                    A1
                         20031117
                                       AU 2003-220480
                                                        20030321
                    A1
                                       EP 2003-716789
    EP 1499319
                B1 20071205
                         20050126
                                                       20030321
    EP 1499319
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                                      AT 2003-716789 20030321
                    T 20071215
    AT 380030
    ES 2297146
                     T3 20080501
                                        ES 2003-716789 20030321
PRIORITY APPLN. INFO.:
                                        US 2002-376731P 20020501
                                        WO 2003-US8893 20030321
GΙ
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GI

10/572,914

AB The title piperazine compds. with general formula of Q-(C=W)m-(CR1R2)n-(C=O)p-T-CO-A [wherein Q = naphthyl, quinolyl, quinoxalinyl, etc.; A = alkoxy, alkyl, cycloalkyl, Ph, or heteroaryl; W = O or NH; T = (un)substituted piperazine; m, n, and p = independently 0-2; R1 and R2 = independently H, OH, alkyl, alkoxy, CN, or F; or R1 and R2 together form CO, CS, C=NH, or (un)substituted C=NOH, etc., with the carbon atom attached] and pharmaceutically acceptable salts thereof are prepared as antiviral agents for the treatment of HIV and AIDS. For example, the compound I was prepared in a multi-step synthesis. I showed EC50 of 0.5 to 5 μ M against human HIV-1 receptors.

Ι

MSTR 1

$$G1 - G2 - G6 - G4$$

$$G1 = 36$$

G8 = 107 / 109

G9—G10 C(O)-G11

G9 = NH

G10 = alkyl <containing 1-6 C>

G11 = NH2

Patent location: claim 1

Note: or pharmaceutically acceptable salts

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 62 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:339137 MARPAT

TITLE: Colorant compositions for light-resistant

high-concentration print images with good color reproducibility and their dispersions, ink-jet inks,

and ink-jet printing process

INVENTOR(S): Takahashi, Mari; Ofuku, Koji; Miura, Norio

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 44 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2003301121	A	20031021	JP 2002-109007	20020411
PRIORITY APPLN. INFO.	:		JP 2002-109007	20020411
GI				

p-C6H4-

AB The compns. contain colorants represented by general formulas selected from (i) I (X1 = II, III, etc.; R1 = H, substituent; m = 0-4 integer; R = substituent; R2, R3 = H, substituent), (ii) X2:N(CR2:CR3)nCR4:Y2 or X3CRa(:CR3CRb)n:NY1 (X2, X3 = coupler residue; R2-R4, Ra, Rb = H, substituent; n = 0, 1, 2; when n = 0, Ra = H, substituent other than electron-withdrawing group; when n = 1, 2, Rb = H, substituent other than electron-withdrawing group; Y1, Y2 = atom group 5- or 6-membered aromatic hydrocarbon ring or heterocyclic ring), or (iii) IV and V (R1 = H, substituent; Y1 = same as above; r = 0, 1, 2, 3). The dispersions contain

VΙ

CO2H

in aqueous media fine particles involving the colorant compns. and polymers and/or high-b.p. organic solvents. The ink-jet inks contain the color compns. or the dispersions. Thus, a water-based colorant composition containing 4%

VI was exemplified.

MSTR 1

G1 = 78

G6 = 100 / CONH2 / SO2NH2

HN----G11

G9 = N G11 = acyl

Patent location: claim 1

Note: additional ring formation also claimed

L5 ANSWER 63 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:307692 MARPAT

TITLE: Preparation of quinoline and related compounds for use

as anti-inflammatory agents

INVENTOR(S): Jaroch, Stefan; Lehmann, Manfred; Schmees, Norbert;

Berger, Markus; Rehwinkel, Hartmut; Krolikiewicz,

Konrad; Skuballa, Werner; Schaecke, Heike;

Schottelius, Arndt

PATENT ASSIGNEE(S): Schering Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 122 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: Ge FAMILY ACC. NUM. COUNT: 1 German

PATENT NO.	KIND I	DATE	APPLICATION NO. DATE
W: AE, CO, HR, LT, PL, UA, RW: GH, KG, FI,	327 A1 2 AG, AL, AM, CR, CU, CZ, HU, ID, IL, LU, LV, MA, PT, RO, RU, UG, UZ, VC, GM, KE, LS, KZ, MD, RU, FR, GB, GR, BJ, CF, CG,	AT, AU, AZ DK, DM, DZ IN, IS, JP MD, MG, MK SC, SD, SE VN, YU, ZA MW, MZ, SD TJ, TM, AT HU, IE, IT	WO 2003-EP3298 20030329 Z, BA, BB, BG, BR, BY, BZ, CA, CH, CN, Z, EC, EE, ES, FI, GB, GD, GE, GH, GM, P, KE, KG, KP, KR, KZ, LC, LK, LR, LS, K, MN, MW, MX, MZ, NI, NO, NZ, OM, PH, E, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, A, ZM, ZW D, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, T, BE, BG, CH, CY, CZ, DE, DK, EE, ES, T, LU, MC, NL, PT, RO, SE, SI, SK, TR, A, GN, GQ, GW, ML, MR, NE, SN, TD, TG DE 2002-10215316 20020402
CA 2481012 AU 20032156 EP 1492771 EP 1492771	A1 2 678 A1 2 A1 2	20031009 20031013	CA 2003-2481012 20030329 AU 2003-215678 20030329 EP 2003-745195 20030329
R: AT, IE, BR 20030089 CN 1659144 JP 20055298 AT 355277 NZ 535872 ES 2282649 US 20040116 US 6897224 TW 272267 MX 2004PA09 NO 2004004 US 20050169 US 7109212 ZA 20040088 US 20060229 US 7329753	BE, CH, DE, SI, LT, LV, A B67 A B61 T A T3 A T3 A B2 B B684 A T31 A B2 B B684 A T31 A B2 B B B B B B B B B B B B B B B B B	DK, ES, FR	AT 2003-745195 20030329 NZ 2003-535872 20030329 ES 2003-745195 20030329 US 2003-405033 20030402 TW 2003-92107522 20030402 MX 2004-PA9684 20041001 NO 2004-4731 20041101 US 2005-59682 20050217 ZA 2004-8827 20060322 US 2006-451508 20060613
PRIORITY APPLN.	INFO.:		DE 2002-10215316 20020402 US 2002-369583P 20020404 WO 2003-EP3298 20030329 US 2003-405033 20030402 US 2005-59682 20050217

AB Title comounds I [A = (un)substituted aryl, benzyl, phenylethyl, etc.; R1, R2 = H, Me, Et, etc.; R3 = alkyl, fluoroalkyl; B = Me or Et substituted methylene, carbonyl; Q = (un)substituted quinoline or isoquinoline] and their pharmaceutically acceptable salts were prepared For example, condensation of 8-quinolinamine and epoxide II afforded quinoline III. Compds. I are noted useful as anti-inflammatory agents (no data provided).

MSTR 1

G16 = quinolinyl (opt. substd. by 1 or more G17) = 147 / 153

G20 = alkylcarbonyl <containing 1-5 C>

Patent location: claim 1

Note: and physiologically acceptable salts

Stereochemistry: and racemates or stereoisomers

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 64 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:265380 MARPAT

TITLE: Hair dye compositions containing quinolinium salts INVENTOR(S): Sauter, Guido; Braun, Hans-Juergen; Duc-Reichlin,

Nadia

10/572,914

PATENT ASSIGNEE(S): Wella Aktiengesellschaft, Germany

SOURCE: Eur. Pat. Appl., 14 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT N	.0	KIND DATE	Ξ	APPLICATION NO.	DATE		
EP 13467	'19	A1 2003	30924	EP 2002-25423	20021115		
R:	AT, BE,	CH, DE, DK,	, ES, FR, GE	s, GR, IT, LI, LU,	NL, SE, MC, PT,		
	IE, SI,	LT, LV, FI,	, RO, MK, CY	, AL, TR, BG, CZ,	EE, SK		
DE 10211	413	A1 2003	30925	DE 2002-10211413	20020315		
US 20030	177592	A1 2003	30925	US 2003-361380	20030210		
US 69770	01	B2 2005	51220				
BR 20030	00496	A 2004	40810	BR 2003-496	20030313		
PRIORITY APPI	N. INFO.	:		DE 2002-10211413	20020315		

AB The invention concerns hair dyes that are prepared from two components; component A1 contains a quinolinium derivative; component A2 includes a nucleophile compound Other direct dyes can be added; solns., emulsions, creams, foams, gels can be formulated. Thus component A1 contained (g): 4-chloro-1-ethylquinolinium tetrafluoroborate 0.70 decyl glycoside 4.0; EDTA disodium salt 0.2; ethanol 5.0; water to 100. Component A2 included: 1,4-diaminobenzene 0.27; decyl glycoside 4.0; EDTA disodium salt 0.2; ethanol 5.0; 25% ammonia solution 6.0; water to 100.

MSTR 2

G2 = CONH2 / 27 / SO2NH2

G7 = NH

Patent location: claim 1

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 65 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:240339 MARPAT

TITLE: Antitumor agent comprising combination of

sulfonamide-containing heterocyclic compound with

angiogenesis inhibitor

INVENTOR(S): Wakabayashi, Toshiaki; Ono, Naoto; Semba, Taro;

Haneda, Toru

PATENT ASSIGNEE(S): Eisai Co., Ltd., Japan SOURCE: PCT Int. Appl., 49 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.					KIND DATE					PPLI	CATI	ои ис	ο.	DATE			
	WO	2003	0740	 45	A1 20030912					M	20	 03-J:	 P249:	2	2003	0304		
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			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
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			UA,	UG,	US,	UΖ,	VC,	VN,	YU,	ZA,	ZM,	ZW						
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			KG,	KΖ,	MD,	RU,	ТJ,	TM,	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
			FΙ,	FR,	GB,	GR,	HU,	ΙE,	ΙT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,
			BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML ,	MR,	ΝE,	SN,	TD,	ΤG
	ΑU	2003	2115	94	A	1	2003	0916		A	U 20	03-2	1159	4	2003	0304		
	EΡ	1481	678		A	1	2004	1201		E:	P 20	03-7	4359	4	2003	0304		
		R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
			ΙE,	SI,	LT,	LV,	FΙ,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK	
	US	2005	0119	303	Α	1	2005	0602		U	S 20	04-5	0467	6	2004	0813		
PRIOR	RIORITY APPLN. INFO.:								J:	P 20	02-5	9471		2002	0305			
										M	0 20	03-J:	P249:	2	2003	0304		
GI																		

AB It is intended to provide compns. and kits for treating tumor whereby the angiogenesis inhibitory activity and the antitumor activity of a sulfonamide-containing heterocyclic compound represented by the following formula (I) can be more effectively exerted. By combining with a VEGF inhibitor or an FGF inhibitor, the sulfonamide-containing heterocyclic compound can be effectively employed in treating cancer.

MSTR 2

G13 = bondG14 = N / 38

с——G15 38

G15 = alkylamino < containing 1-4 C>

(opt. substd. by 1 or more G2) / CONH2

 $G16 = 42-5 \ 43-8$



Patent location: claim 7

Note: substitution is restricted

Note: additional ring formation also claimed

REFERENCE COUNT: 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 66 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:214614 MARPAT

TITLE: Preparation of N-(azabicyclyl)arylamides for

therapeutic use as nicotinic acetylcholine receptor

agonists

INVENTOR(S): Jacobsen, Eric Jon; Myers, Jason K.; Walker, Daniel

P.; Wishka, Donn G.; Reitz, Steven C.; Piotrowski, David W.; Acker, Brad A.; Groppi, Vincent E., Jr.

PATENT ASSIGNEE(S): Pharmacia & Upjohn Company, USA

SOURCE: PCT Int. Appl., 145 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

FAMILI ACC. NUM. COUNT:

PATENT NO. KIND						DATE			A	PPLI	CATI	ON No	Ο.	DATE				
									_									
WO	O 2003072578 A1					2003	0904		M	0 2 0	03-U	S268	8	20030214				
	W:	ΑE,	AG,	AL,	ΑM,	ΑT,	ΑU,	ΑZ,	ΒA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,	
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,	
		LS.	LT.	LU.	LV.	MA.	MD.	MG.	MK.	MN.	MW.	MX.	MZ.	NO.	NZ.	OM.	PH.	

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PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ,
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             KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     CA 2475773
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                           20030904
                                           CA 2003-2475773 20030214
     AU 2003214936
                       Α1
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                                           AU 2003-214936
     US 20030236270
                            20031225
                                           US 2003-366894
                                                             20030214
                       Α1
     US 7001900
                            20060221
                       В2
     EP 1478646
                       Α1
                            20041124
                                           EP 2003-710784
                                                             20030214
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             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
     BR 2003007874
                            20041228
                                           BR 2003-7874
                                                             20030214
                      Α
     JP 2005525357
                                           JP 2003-571284
                       Τ
                            20050825
                                                             20030214
     MX 2004PA07083
                                           MX 2004-PA7083
                            20041029
                                                             20040722
                       Α
PRIORITY APPLN. INFO.:
                                           US 2002-358146P 20020220
                                           WO 2003-US2688
                                                            20030214
GΙ
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AB N-(azabicyclyl) arylamides, such as RNR1C(:X)W [R = azabicyclyl; R1 = H, alkyl, cycloalkyl, haloalkyl, aryl; W = heteroaryl; X = O, S], were prepared for therapeutic use as nicotinic acetylcholine receptor agonists. These amides are useful for the treatment of central nervous system disorders, such as cognitive and attention deficit symptoms of Alzheimer's, neurodegeneration associated with diseases such as Alzheimer's disease, pre-senile dementia (mild cognitive impairment), senile dementia, schizophrenia, psychosis, attention deficit disorder, attention deficit hyperactivity disorder, mood and affective disorders, amyotrophic lateral sclerosis, borderline personality disorder, traumatic brain injury, behavioral and cognitive problems associated with brain tumors, AIDS dementia complex, dementia associated with Down's syndrome, dementia associated with

Lewy

Bodies, Huntington's disease, depression, general anxiety disorder, age-related macular degeneration, Parkinson's disease, tardive dyskinesia, Pick's disease, post traumatic stress disorder, dysregulation of food intake including bulimia and anorexia nervosa, withdrawal symptoms associated with smoking cessation and dependent drug cessation, Gilles de la Tourette's Syndrome, glaucoma, neurodegeneration associated with glaucoma, or symptoms associated with pain. Thus, the hydrochloride salt of amide I was prepared via a multistep synthetic sequence which concluded with an amidation reaction of the corresponding (2S,3R)-azabicyclic amine dihydrochloride with 2-naphthoic acid using diphenylphosphinic chloride and Et3N in THF. The prepared amides were assayed for human $\alpha7-5\mathrm{HT3}$

receptor binding activity.

MSTR 1

G6 = 191

G19 = 81 / 136

G22-G23 C(0)-G24

G22 = NH

G23 = alkyl < containing 1-4 C >

(opt. substd. by 1 or more G12)

G24 = NH2

Patent location: claim 1

Note: substitution is restricted

Note: or pharmaceutically compositions or pharmaceutically acceptable salts

Note: or racemic mixtures or pure enantiomers

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 67 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:180085 MARPAT

TITLE: Preparation of novel aryl- and heteroarylpiperazines

with histamine H3 receptor affinity

INVENTOR(S): Hohlweg, Rolf; Doerwald, Florencio Zaragoza;

Stephensen, Henrik; Pettersson, Ingrid; Peschke, Bernd

PATENT ASSIGNEE(S): Novo Nordisk A/S, Den.; Boehringer Ingelheim

International G.m.b.H.
PCT Int Appl 145 pp

SOURCE: PCT Int. Appl., 145 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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A2
                            20030814
                                           WO 2003-DK71
     WO 2003066604
                                                             20030205
     WO 2003066604
                            20031204
                      А3
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
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             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     CA 2474214
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                       Α2
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                            20041110
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         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
     BR 2003007429
                            20041228
                                           BR 2003-7429
                                                             20030205
                     Α
     CN 1628109
                       Α
                            20050615
                                           CN 2003-803360
                                                             20030205
                                           JP 2003-565978
     JP 2005533747
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                                                             20030205
                                           US 2003-383310
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                                           ZA 2004-5694
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                       Α
                                                             20040903
PRIORITY APPLN. INFO.:
                                           DK 2002-168
                                                             20020205
                                           US 2002-356630P
                                                             20020208
                                           DK 2002-1142
                                                             20020726
                                           US 2002-399304P
                                                            20020726
                                           WO 2003-DK71
                                                             20030205
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GΙ

AB Novel aryl- and heteroarylpiperazines of formula I [R1 = alkyl, alkenyl, alkynyl, cycloalkyl, not isobutyl; R2 = H, alkyl; R1R2 = alkylene; R3 = H, halo, OH, CF3, OCF3, alkyl, cycloalkyl, alkoxy, aryl, etc.; A = aryl, heteroaryl, etc.] are prepared and used in pharmaceutical compns. The compds. show a high and selective binding affinity to the histamine H3 receptor indicating histamine H3 receptor antagonistic, inverse agonistic or agonistic activity. As a result, the compds. are useful for the treatment of diseases and disorders related to the histamine H3 receptor. Thus, II was prepared from 1-(4-hydroxyphenyl)piperazine and cyclopentanone in 49% yield.

ΙI

MSTR 1

G3 = 53

G7 = 95 / alkylamino <containing 1-6 C> (opt. substd.)

95 (O)-G13

G13 = NH2 / heterocycle <containing 1 heteroatom, 1 N,

3-6 C, attached through 1 N, monocyclic>

Patent location: claim 1

Note: substitution is restricted

Note: additional ring formation also claimed

Note: also incorporates claim 57

L5 ANSWER 68 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:143997 MARPAT

TITLE: Methods using Edg receptor modulators for the treatment of Edg receptor-associated conditions

INVENTOR(S): Shankar, Geetha; Solow-Cordero, David; Spencer, Juliet

V.; Gluchowski, Charles

PATENT ASSIGNEE(S): Ceretek LLC, USA SOURCE: PCT Int. Appl., 293 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PATENT NO.	KIND DAT	TE AI	PPLICATION NO.	DATE
WO 2003062392	A2 200		O 2003-US1881	20030121
WO 2003062392	A3 200	050120		
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CO, CR,	CU, CZ, DE	E, DK, DM, DZ,	EC, EE, ES, FI,	GB, GD, GE, GH,
GM, HR,	HU, ID, IL	L, IN, IS, JP,	KE, KG, KP, KR,	KZ, LC, LK, LR,
LS, LT,	LU, LV, MA	A, MD, MG, MK,	MN, MW, MX, MZ,	NO, NZ, OM, PH,

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PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
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                          20030731
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     EP 1513522
                      A2
                            20050316
                                         EP 2003-710713
                                                            20030121
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                                          JP 2003-562260
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PRIORITY APPLN. INFO.:
                                           US 2002-350445P 20020118
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                                           US 2002-350447P
                                                           20020118
                                           US 2002-350448P 20020118
                                           WO 2003-US1881
                                                            20030121
                                           US 2003-352579
                                                           20030127
```

AB The invention provides a method of modulating an Edg-2, Edg-3, Ed-4 or Edg7 receptor-mediated biol. activity in a cell. A cell expressing the Edg-2, Edg-3, Edg-4 or Edg 7 receptor is contacted with a modulator of the Edg-2, Edg-3, Ed-4 or Edg 7 receptor sufficient to modulate receptor mediated biol. activity. In another aspect, the present invention provides a method for modulating an Edg-2, Edg-3, Ed-4 or Edg-7 receptor mediated biol. in a subject. A therapeutically effective amount of a modulator of the Edg-2, Edg-3, Ed-4 or Edg7 receptor is administered to the subject. Preparation of compds., e.g.

4,4,4-trifluoro-3-oxo-N-(5-phenyl-2H-pyrazol-3-yl)butyramide, is described.

MSTR 20

G1 = 34-5 35-2

G2 = 63 / CONH2 (opt. substd.)

G5 = CONH2 (opt. substd.)

Patent location: claim 135

Note: or pharmaceutically available solvates or hydrates

L5 ANSWER 69 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:69267 MARPAT

TITLE: Preparation of 2-benzimidazolylamines as ORL1-receptor

agonists for the treatment of pain and inflammatory

diseases

SOURCE: Eur. Pat. Appl., 33 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

-	PAI	ENT	NO.		KII	ND	DATE			AP	PLI	ICATI	ON NO).	DATE			
	 EP	1069	 124		 A:	1	2001	0117		EP	20	 000-3	05981	 L	20000	714		
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		R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
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1	US	6340	681		В	1	2002	0122		US	2(000-6	06923	l	20000	629		
	JΡ	2001	0488	79	А		2001	0220		JP	2(000-2	09374	1	20000	711		
	JΡ	3276	111		Βź	2	2002	0422										
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	PΤ	1069	124		T		2004	0930		PT	20	000-3	05983	L	20000	714		
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(CA	2314	800		A.	1	2001	0116		CA	. 20	000-2	31400	8 (20000	717		
PRIOR	ΙTΊ	APP	LN.	INFO	. :					WC	19	999-I	B1290)	19990	716		
GI																		

AB Title compds. I [R1, R2 = H, halo, OH, etc.; R3, R4 = H, halo-alkyl, substituted alkyl, i.e., OH, alkoxy, alkyl-S, etc.; R5 = pheny, substituted cycloalkyl, i.e., H, halo, OH, etc.;] and their pharmaceutically acceptable salts were prepared For example, N-alkylation of N-methylpiperazine by chlorobenzimidazolyl II, e.g., prepared from 1,3-dihydro-1-(4-piperidinyl)-2H-benzimidazol-2-one in 2-steps, afforded 2-benzimidazolylamine III in 15% yield. In selective affinity studies of opioid receptors, i.e., ORL1, $\mu, \, \kappa$ and δ , some examples of compds. I exhibited good ORL1-receptor agonist activity. Compds. I are claimed useful as analgesics.

MSTR 1

^{*} STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

$$G1$$
 $G1$ $G1$ N $G3$ N $G5$

G3 = 348

G36 = 473 / alkoxycarbonylamino <containing 1-4 C>

C(O)-G45

G45 = NH2

Patent location: claim 1 Note: or salts

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 70 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:53194 MARPAT

TITLE: Preparation of bicyclic N-arylamides for use in

producing pharmaceuticals

INVENTOR(S): Luithle, Joachim; Boess, Frank-Gerhard; Erb,

Christina; Flessner, Timo; Hendrix, Martin; Van

Kampen, Marja; Methfessel, Christoph
: Bayer Aktiengesellschaft, Germany

PATENT ASSIGNEE(S): Bayer Aktiengesellschaft SOURCE: PCT Int. Appl., 36 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT N	.0		KII	ND	DATE			A)	PPLI	CATI	и ис	Э.	DATE			
WO 20030	51874	4	A.	1	2003	0626		M	20	02-E	P138	35	2002	1206		
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	GM, F	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
	LS, I	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NO,	NΖ,	OM,	PH,
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     US 20050107460
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PRIORITY APPLN. INFO.:
                                            DE 2001-10162375 20011219
                                            WO 2002-EP13835 20021206
OTHER SOURCE(S):
                        CASREACT 139:53194
GΙ
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The invention relates to novel bicyclic N-arylamides, R1C(:0)NR2R3 [R1 = 1-azabicyclo[m.n.p]alkyl (7 - 11 ring atoms, optionally substituted with C1-6-alkyl); m, n = 2, 3; p = 1, 2, 3; R2 = 8 - 10 membered heteroaryl, naphthyl, azulenyl (optionally substituted with H, halogen, CHO, CONH2, CN, CF3, CF3O, NO2, C1-6-alkyl,C1-6-alkoxy, C1-6-alkylthio); R3 = H, C1-6-alkyl] and their salts, solvates and salt solvates, to a method for the production thereof, characterized by reaction of R1COX [X = OH, appropriate leaving group] with R2R3NH in the presence of a base, and to the use of the same for producing pharmaceuticals for the treatment and/or prophylaxis of diseases and for improving perception, power of concentration, learning capacity and/or memory retention. Thus, N-(6-quinoxalinyl)quinuclidine-3-carboxamide hydrochloride (I·HCl) was prepared from quinuclidine-3-carboxamide hydrochloride and (6-quinoxalinyl)amine in DMF containing EtN(CHMe2)2 and catalytic DMAP.

MSTR 1

G7---G4

G2 = NH G4 = quinolinyl (opt. substd. by 1 or more G5) 10/572,914

G5 = CONH2 G7 = 3



Patent location: claim 1

Note: and salts, solvates and solvates of salts

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 71 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:36536 MARPAT

TITLE: Preparation of quinoline and quinazoline derivatives

as inflammation modulators

INVENTOR(S): Cushing, Timothy D.; He, Xiao; Smith, Marie-Louise;

Degraffenreid, Michael R.; Powers, Jay; Tomooka, Craig S.; Clark, David L.; Hao, Xiaolin; Jaen, Juan C.;

Labelle, Marc; Walker, Nigel P. C.; Gill, Adrian L.;

Talamas, Francisco X.; Labadie, Sharada S. Tularik Inc., USA; F. Hoffmann-La Roche AG

PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 102 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

	PAT	PATENT NO.				ND	DATE			APPLICATION NO. DATE										
		2003048152								W	0 20	02-U	20021204							
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		W:	ΑE,	ΑG,	ΑL,	ΑM,	ΑT,	ΑU,	ΑZ,	ΒA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,		
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			GM.	HR.	HU.	ID.	IL,	IN.	IS.	JP.	KE.	KG.	KP.	KR.	KZ.	LC.	LK.	LR.		
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		PL, PT,															•			
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		D TaT a	,	,	,	,	,	,	,	•	•		TIC	71/1	17 TaT	7\ 1\/I	7\ '7	DV		
		KW:					MW,													
			•	•	•	•	ТJ,	•	•	•	•	•	•	•	•	•	•	•		
							IE,	•		•	•			•			BF,	ВJ,		
			CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML ,	MR,	ΝE,	SN,	TD,	ΤG				
	AU 2002365611					A1 20030617				A.	U 20	02-3	1	20021204						
	US 20030181472					A1 20030925				U	S 20	02-3	8	20021204						
	US 7176314						B2 20070213													
PRIO	PRIORITY APPLN. INFO.						. •			US 2001-337460P						20011205				
															20021204					
GI										• • • • • • • • • • • • • • • • • • • •	0 20	01 0		-		01				
\circ																				

AB Title compds. I [X = N, (un)substituted CH; J = alkylene, alkenylene, alkynylene, CO, C:S, (un)substituted C:NH, NH, CONH, CSNH, C(:NH)NH, CH:N, O, S, S(O), SO2, alkylenamino, alkylenoxy; K = bond, alkylene, CO, CS, O, S, S(O), SO2, (un)substituted C:NH, NH; L = H, (un)substituted OH, alkyl, heteroalkyl, aryl, heteroaryl, NH2, acyl, thioacyl, CH:NH, carbamoyl, thiocarbamoyl, CO2H; JK, JL, KL = heterocyclic; B = 5-6-membered heteroarom.; R, R1 = H, halogen, (un)substituted alkyl, alkenyl, alkynyl, alkoxy, alkylthio, NH2, cycloalkyl, heterocyclic, CN, NO2, acyl, alkoxycarbonyl, CONH2, SO2NH2] were prepared for use in the treatment of inflammatory, immunoregulatory, metabolic and cell proliferative conditions or diseases. Thus, 5-chloroisatin was iodinated, cyclized with 5-acetyl-1-methyl-2-tert.-butyldimethylsilylimdazole, substituted with CH2:CHCN, reduced, and treated with 4-methylpyridine to give the quinoline II. I had IC50 ≤ 30 μM for inhibition of IKKβ.

MSTR 1

= CONH2

G5

G10 = SO2NH2

G12 = NH (opt. substd.)

G13 = Ph

Patent location: claim 1

Note: or pharmaceutically acceptable salts or prodrugs

Note: substitution is restricted

ANSWER 72 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

139:36445 MARPAT ACCESSION NUMBER:

TITLE: Preparation of 2-aminoquinolines as melanin

concentrating hormone receptor (MCH-1R) antagonists. INVENTOR(S): Devita, Robert J.; Chang, Lehua; Chaung, Danny; Hoang,

Myle; Jiang, Jinlong; Lin, Peter; Sailer, Andreas W.;

Young, Jonathan R.

PATENT ASSIGNEE(S): Merck & Co., Inc., USA PCT Int. Appl., 178 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

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PATENT NO. KIND DATE
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    _____
    WO 2003045313 A2 20030605
WO 2003045313 A3 20030904
                                        WO 2002-US37556 20021122
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            LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL,
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                    A1 20030605 CA 2002-2468015 20021122
    AU 2002352878
                    A1 20030610
                                       AU 2002-352878 20021122
    AU 2002352878
                    В2
                         20071122
    EP 1450801
                    A2 20040901
                                       EP 2002-789837 20021122
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    JP 2005519876 T 20050707
                                       JP 2003-546818 20021122
    US 20050026915
                    A1 20050203
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                                                       20040525
    US 7084156 B2 20060801
PRIORITY APPLN. INFO.:
                                        US 2001-333581P 20011127
                                        WO 2002-US37556 20021122
GΙ
```

AB Title compds. [I; R1, R2 = H, (substituted) alkyl, alkenyl, alkynyl, cycloalkylalkyl, aralkyl, etc.; R1R2N = 4-11 membered (bridged) (substituted) heterocyclyl; R3, R4 = H, halo, (substituted) alkyl, alkenyl, alkynyl, perfluoroalkyl, cycloalkyl, cycloalkylalkyl, aryl, aralkyl, heteroaralkyl, OR7, N(R7)2, cyano, etc.; R3R4 = atoms to form 5-7 membered (substituted) ring; R5 = H, halo, alkyl, perfluoroalkyl, OR7, N(R7)2; R6 = (CH2)nR7, (CH2)nCN, (CH2)nCO2R7, (CH2)nOR7, (CH2)nN(R7)2, etc.; R7 = H, alkyl, aryl, heteroaryl, cycloalkyl, aralkyl, aralkenyl, cycloalkylalkenyl, etc.; n = 0-5], were prepared for the treatment or prevention of obesity, eating disorders, osteoarthritis, cancer, AIDS wasting, cachexia, frailty, mental disorders, stress, cognitive disorders, sexual function, reproductive function, kidney function, locomotor disorders, attention deficit disorder (ADD), substance abuse disorders and dyskinesias, Huntington's disease, epilepsy, memory function, and spinal muscular atrophy. Thus, 2-piperidin-1-ylquinolin-6-amine and (2E)-3-(4-chlorophenyl)prop-2-enoyl chloride were stirred 3 h in HOAc to give (2E)-3-(4-chlorophenyl)-N-(2-piperidin-1-ylquinolin-6-yl)prop-2enamide hydrochloride. I bound to MCH-1R receptors with IC50 = 0.1-10000nM.

MSTR 1

G9 = 48

G11 = heterocycle <containing 3 or more atoms,
 zero or more N, zero or more O,
 zero or more S (no other heteroatoms),
 0 or more double bonds, mono- or polycyclic,
 including 5- or 6-membered rings> (opt. substd.) / Ph
G15 = 107 / 136

G18-G11 G17-G19

G18 = S02G19 = 173

173 G18 G11

G21 = 261

261 G11

Patent location: claim 1

Note: and pharmaceutically acceptable salts

Note: substitution is restricted

Note: additional substitution also claimed

L5 ANSWER 73 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 139:22115 MARPAT

TITLE: Preparation of 4-aminoquinolines as melanin

concentrating hormone receptor antagonists,

particularly MCH-1R antagonists.

INVENTOR(S): Devita, Robert J.; Chang, Lehua; Hoang, Myle Thi;

Jiang, Jinlong; Lin, Peter; Sailer, Andreas W.

PATENT ASSIGNEE(S): Merck & Co., Inc., USA SOURCE: PCT Int. Appl., 159 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PAT	FENT	NO.		KIND		DATE			A	PPLI	CATI	N NC	Ο.	DATE			
WO 2003045920				A	1	2003	0605		W	0 20	1122						
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		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KR,	KΖ,	LC,	LK,	LR,	LS,
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GΙ

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, SK
JP 2005518365 T 20050623 JP 2003-547372 20021122
US 20050009815 A1 20050113 US 2004-496614 20040525
PRIORITY APPLN. INFO:: US 2001-333464P 20011127
WO 2002-US37510 20021122

AΒ Title compds. [I; R1 R2 = H, (substituted) alkyl, alkenyl, alkynyl, cycloalkylalkyl, heterocycloalkylalkyl, arylalkyl, heteroarylalkyl; R1R2N = (substituted) heterocyclyl; R3, R4 = H, halo, (substituted) alkyl, perfluoroalkyl, alkenyl, alkynyl, cycloalkyl, cycloalkylalkyl, cycloheteroalkyl, cycloheteroalkylalkyl, aryl, arylalkyl, heteroaryl, heteroarylalkyl, OR7, NR7R7, CO2R7, cyano, CONR7R7; R3R4 = atoms to form a (substituted) 5-7 membered (hetero)cycloalkyl; R5 = H, halo, alkyl, perfluoroalkyl, OR7, NR7R7; R6 = (CH2)nR7, (CH2)naryl-R7, (CH2)n-heteroaryl-R7, (CH2)n-heterocycloalkyl-R7, (CH2)nCN, (CH2) nCON(R7)2, (CH2) nCO2R7, (CH2) nCOR7, (CH2) nNR7COR7, (CH2) nNR7CO(CH2) nSR7 (CH2) nNR7CO2R7, (CH2) nNR7CON(R7)2, (CH2) nNR7SO2R7, (CH2)nSOpR7, (CH2)nSO2N(R7)2, (CH2)nOR7, (CH2)nOC(O)R7, (CH2)nOCO2R7, (CH2)nO2CN(R7)2, (CH2)nN(R7)2, (CH2)nNR7SO2N(R7)2; R7 = H, (substituted) alkyl, aryl, heteroaryl, cycloalkyl, heterocycloalkyl, aralkyl, heteroarylalkyl, cycloalkylalkyl, heterocycloalkylalkyl, arylalkenyl, heteroarylalkenyl, cycloalkylalkenyl, heterocycloalkylalkenyl; n = 0-5; p = 0-2], were prepared Thus, 2-propylquinoline-4,6-diamine and (2E)-3-(4-chlorophenyl)prop-2-enoyl chloride were stirred 6 h in HOAc to qive (2E)-N-(4-amino-2-propylquinolin-6-yl)-3-(4-chlorophenyl)prop-2enamide. I are useful for the treatment or prevention of obesity or eating disorders, osteoarthritis, certain cancers, AIDS wasting, cachexia, frailty, mental disorders, stress, cognitive disorders, sexual function, reproductive function, kidney function, locomotor disorders, attention deficit disorder, substance abuse disorders, dyskinesias, Huntington's disease, epilepsy, memory function, and spinal muscular atrophy. I showed IC50 = 0.1-10000 nM for MCH-1R receptor binding activity.

MSTR 1

10/572,914

G1 = 14

M 14 G2

G2 = cyclopropyl

G9 = 48

G11 = heterocycle <containing 3 or more atoms,

zero or more N, zero or more O,

zero or more S (no other heteroatoms),

O or more double bonds, mono- or polycyclic, including 5- or 6-membered rings> (opt. substd.)

G15 = 107 / 136

G18—G11 G17—G19

G18 = S02G19 = 173

G18-G11

Patent location: claim 1

Note: and pharmaceutically acceptable salts

Note: substitution is restricted

Note: additional substitution also claimed

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 74 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 138:255107 MARPAT

TITLE: Synthesis of enantiomerically pure amino-substituted

fused bicyclic rings

INVENTOR(S): McEachern, Ernest J.; Bridger, Gary J.; Skupinska,

Krystyna A.; Skerlj, Renato T.

PATENT ASSIGNEE(S): Anormed Inc., Can. SOURCE: PCT Int. Appl., 85 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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                                           WO 2002-US29372 20020912
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PRIORITY APPLN. INFO.:
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                                           WO 2002-US29372 20020912
                                           US 2004-959823
                                                            20041006
GΙ
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AB This invention describes various processes for synthesis and resolution of racemic amino-substituted fused bicyclic ring systems (shown as I; variables defined below), primarily 5,6,7,8-tetrahydroquinolines (shown as II; e.g. 8-amino-2-methyl-5,6,7,8-tetrahydroquinoline) and 5,6,7,8-tetrahydroisoquinolines (e.g. 5-amino-5,6,7,8-tetrahydroisoquinoline). One process uses selective hydrogenation of an amino-substituted fused bicyclic aromatic ring system; for example,

8-amino-5,6,7,8-tetrahydroquinoline was obtained in 2 steps (100, 54 and 91% yields) from 8-aminoquinoline via intermediates 8-acetylaminoquinoline and 8-acetylamino-5,6,7,8-tetrahydroquinoline using PtO2/trifluoroacetic acid/H2 for the hydrogenation. An alternative process preps. the racemic amino-substituted fused bicyclic ring system via nitrosation; for example, 5,6,7,8-tetrahydroquinoline was converted with LDA/MTBE at -30° followed by isoamyl nitrite to 8-hydroxyimino-5,6,7,8-tetrahydroquinoline (75%) that was hydrogenated using H2/Pd/C/MeOH to give 8-amino-5,6,7,8-tetrahydroquinoline (100%). The present invention describes the enzymic resolution of a racemic mixture to produce the (R)- and (S) - forms of amino-substituted fused bicyclic rings as well as a racemization process to recycle the unpreferred enantiomer. For example, 8-amino-5,6,7,8-tetrahydroquinoline was half reacted with EtOAc in iPr20 at 60° in the presence of Candida antarctica lipase to give (R)-(-)-N-(5,6,7,8-tetrahydroquinolin-8-yl) acetamide (97% ee) and unreacted (S)-(+)-8-amino-5,6,7,8-tetrahydroquinoline (96% ee). The amine could be racemized at 150° in a sealed tube under Ar with 87% yield. Further provided by this invention is an asym. synthesis of the (R) - or (S) - enantiomer of primary amino-substituted fused bicyclic ring systems. For example, (R)-(-)-8-amino-5,6,7,8-tetrahydroquinoline (98%)ee) was obtained in 4 steps (82, 95, 93 and 59% yields) starting from 8-hydroxy-5,6,7,8-tetrahydroquinoline via intermediates 6,7-dihydro-5H-quinolin-8-one, (R)-(-)-(6,7-dihydro-5H-quinolin-8-one)ylidene)(1-phenylethyl)amine, and (-)-((1R)-1-Phenylethyl)-(8-(R)-5,6,7,8-1)tetrahydroquinolin-8-yl)amine using $(R)-(+)-\alpha$ -methylbenzylamine as chiral auxiliary. For I: ring A is a heteroarom. 5- or 6-membered ring, P is N, S or O; ring B is a 5- or 6-membered cycloalkyl or heterocycloalkyl; NH2 is located at a position on ring B; and R2 is located at any other H position on the fused bicyclic ring; m is 0-4; R2 = halo, nitro, cyano, carboxylic acid, alkyl, alkenyl, cycloalkyl, hydroxy, thiol, a protected amino, acyl, carboxylate, carboxamide, sulfonamide, an aromatic group and a heterocyclic group. The variable definitions for II and the isoquinolines are the same as for I.

MSTR 1

G1-NH₂

G1 = 5

G2 = 140 / CONH2 (opt. substd.) / SO2NH2 (opt. substd.)

HN----G3

G3 = acyl

Patent location: claim 1

Note: substitution is restricted

L5 ANSWER 75 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 138:180679 MARPAT

TITLE: SH3 protein domains and their ligands

INVENTOR(S): Booker, Grant William; Pyke, Simon Mathew; Branson,

Kim Mathew; Inglis, Steven Robert

PATENT ASSIGNEE(S): Adelaide Research & Innovation Pty Ltd., Australia

SOURCE: PCT Int. Appl., 176 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	CENT 1	NO.		KIND DATE					A.	PPLI	CATI	DATE					
	WO	WO 2003013523			A1 20030220				M.	0 20	 02-A	J106	4	20020808				
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			CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,
			PT,	SE,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,
			ΝE,	SN,	TD,	ΤG												
AU 2002319011 A1 20030224										AU 2002-319011 20020808								
PRIORITY APPLN. INFO.:									AU 2001-6881 20010808									
										WO 2002-AU1064 20020808								

The present invention relates generally to mols. capable of interaction AΒ with one or more domains within a proteinaceous mol. such as a peptide, polypeptide, protein or a macromol. comprising a proteinaceous mol. More particularly the present invention relates to mols. including ligands which are capable of interacting with, and more particularly, binding to, SH3 protein domains or homologs thereof and even more particularly to mols. including ligands which are capable of binding to SH3 domains having a three-dimensional ligand-binding site comprising a neg. charged residue and a hydrophobic residue linearly separated by at least five amino acid residues. The subject invention is preferably directed to the use of 2-aminopyridine, 2-aminoquinoline, 1-aminoisoquinoline and derivs., homologs, analogs and mimetics thereof or pharmaceutically acceptable salts thereof which interact with SH3 domains, and more particularly to the binding of 2-aminopyridine, 2-aminoquinoline, 1-aminoisoquinoline and derivs. analogs and mimetics to SH3 domains as defined above. The present invention contemplates the use of a three dimensional structure of the subject SH3 domain to identify, screen and design amino-substituted and amino-substituted pyridines and aminoquinolines capable of binding to an SH3 domain. The present invention is also useful for the in silico selection of derivs. homologs, analogs and mimetics of 2-aminopyridine, 2-aminoquinoline, 1-aminoisoquinoline capable of binding to SH3 domains. The ligands of the present invention are useful in the development of a

range of therapeutic and diagnostic agents.

MSTR 2

G1---G5

G1 = 23

G2 = CONH2 / alkylamino <containing 1-12 C>

(opt. substd.)

Patent location: claim 1

Note: additional oxo group substitution, fused ring

formation, and unsaturation also claimed

Note: substitution is restricted

Note: or pharmaceutically acceptable salts or other

derivatives

Stereochemistry: or diastereoisomers

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d ibib abs fqhit 76-125

L5 ANSWER 76 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 137:325443 MARPAT

TITLE: Preparation of novel tricyclic benzodiazepine

carboxamides as tocolytic oxytocin receptor

antagonists

INVENTOR(S): Failli, Amedeo Arturo; Shumsky, Jay Scott; Caggiano,

Thomas Joseph; Sabatucci, Joseph Peter; Memoli, Kevin

Anthony; Trybulski, Eugene John

PATENT ASSIGNEE(S): Wyeth, John, and Brother Ltd., USA

SOURCE: PCT Int. Appl., 158 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE
WO 2002083683 A1 20021024 WO 2002-US11534 20020411
WO 2002083683 A9 20040226

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,

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            PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
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    EP 1377581
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PRIORITY APPLN. INFO.:
                                          US 2001-283262P 20010412
                                          WO 2002-US11534 20020411
GΙ
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The title compds. [I; ring containing Z = II, III (wherein R1, R2 = H, alkyl, AB halo, etc.); R3 = H, alkyl, alkoxy, etc.; R4 = BC (B = IV, V; C = (un) substituted Ph, 1-naphthyl, 1-pyrrolyl, etc.; A = CH, N; R5-R7 = H, alkyl, alkoxy, etc.); R = pyridylmethylamino, 2-(pyridyl)ethylamino, 4-(pyridyl)piperazino, etc.] which act as oxytocin receptor competitive antagonists, and therefore are useful in treatment, inhibition, suppression or prevention of preterm labor, dysmenorrhea, endometritis, suppression of labor at term prior to Caesarian delivery, and to facilitate antinatal transport to a medical facility, were prepared Thus, a 7-step synthesis of VI which showed IC50 of 11.2 nM against human oxytocin receptor binding, was given. The compds. I are also useful in enhancing fertility rates, enhancing survival rates and synchronizing estrus in farm animals, and may be useful in the prevention and treatment of dysfunctions of the oxytocin system in the central nervous system including obsessive compulsive disorder (OCD) and neuropsychiatric disorders.

MSTR 1

G5 = 417

G6 = N

G7 = CONH2 / alkylamino <containing 1-6 C> / SO2NH2

Patent location: claim 1

Note: or pharmaceutically acceptable salts

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 77 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 137:325440 MARPAT

TITLE: Preparation of novel tricyclic benzodiazepine

carboxamides as tocolytic oxytocin receptor

antagonists

INVENTOR(S): Failli, Amedeo Arturo; Shumsky, Jay Scott; Caggiano,

Thomas Joseph; Sabatucci, Joseph Peter; Memoli, Kevin

Anthony; Trybulski, Eugene John; Sanders, William

Jennings

PATENT ASSIGNEE(S): Wyeth, John, and Brother Ltd., USA

SOURCE: PCT Int. Appl., 149 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PRIORITY APPLN. INFO.:
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                                          WO 2002-US11530 20020411
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The title compds. [I; ring containing Z = II, III (wherein R1, R2 = H, alkyl, AB halo, etc.); R3 = H, alkyl, alkoxy, etc.; R4 = BC (B = IV, V; C =(un) substituted Ph, 1-naphthyl, 1-pyrrolyl, etc.; A = CH, N; R5-R7 = H, alkyl, alkoxy, etc.)] which act as oxytocin receptor competitive antagonists, and therefore are useful in treatment, inhibition, suppression or prevention of preterm labor, dysmenorrhea and endometritis, suppression of labor at term prior to Caesarian delivery, and to facilitate antinatal transport to a medical facility, were prepared E.g., a 7-step synthesis of VI which showed IC50 of 1.37 nM against human oxytocin receptor binding (CHO cell line), was given. The compds. I are also useful in enhancing fertility rates, enhancing survival rates and synchronizing estrus in farm animals, and may be useful in the prevention and treatment of disfunctions of the oxytocin system in the central nervous system including obsessive compulsive disorder (OCD) and neuropsychiatric disorders.

MSTR 1

G6 = N

G7 = CONH2 / alkylamino <containing 1-6 C> / SO2NH2

Patent location: claim 1

Note: and pharmaceutically acceptable salts, or prodrugs

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 78 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 137:310939 MARPAT

TITLE: Preparation of tricyclic diazepines as tocolytic

oxytocin receptor antagonists

INVENTOR(S): Failli, Amedeo Arturo; Shumsky, Jay Scott; Caggiano,

Thomas Joseph; Sabatucci, Joseph Peter; Memoli, Kevin

Anthony; Trybulski, Eugene John

PATENT ASSIGNEE(S): Wyeth, John and Brother Ltd., USA

SOURCE: PCT Int. Appl., 220 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA7	CENT	NO.		KI	ND	DATE APPLICATION NO. D.				DATE							
WO	2002	0836	78	А	1	2002	1024		M	O 20	02-U	S115	27	2002	0411		
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		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NΖ,	OM,	PH,
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	UZ,	VN,	YU,	ZA,	ZM,	ZW								
	RW:													ZW,			
														NL,			
														ΝE,		TD,	ΤG
									U	S 20	02-1	1997	1	2002	0410		
	7109																
														2002			
	2002									-	-		_	2002	-		
	1377								E.	P 20	02 - 7	3134	3	2002	0411		
EP	1377																
	R:											LI,	LU,	NL,	SE,	MC,	PT,
		•				FI,							_				
	1501													2002			
	2004										02-5			2002			
	2002													2002			
														2002			
ES	2260	434		Т	3	2006	1101		E	S 20	02-7.	3134	3	2002	0411		

MX 2003PA09331 A 20041112 PRIORITY APPLN. INFO.:

MX 2003-PA9331 20031010 US 2001-283264P 20010412 WO 2002-US11527 20020411

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AΒ The title compds. [I; ring containing Z = II, III; R1, R2 = H, alkyl, halo, CN, etc.; R3 = H, alkyl, alkoxy, etc.; R4 = BC (wherein B = IV, V; C = (un) substituted Ph, 1-naphthyl, 1-pyrrolyl, etc.; A = CH, N; R5-R7 = H, alkyl, alkoxy, etc.); R = OH, NR11R12, (un)substituted 4-oxopiperidin-1-yl, etc. (R11, R12 = H, alkyl, cycloalkyl, etc.)], useful for the treatment and/or prevention and/or suppression of disorders which may be remedied or alleviated by oxytocin antagonist activity, including treatment of preterm labor, dysmerorrhea, endometritis, and for suppressing labor prior to Caesarian delivery, were prepared Thus, amidation of VI [R = OH] (multi-step synthesis given) with 1-(tert-butoxycarbonyl)piperazine afforded VI [R = 4-tertbutoxycarbonylpiperazin-1-yl] which showed 56% inhibition of binding to membranes of CHO cell line stably transfected with human oxytocin receptor at 100 nM vs. 2% and 13% inhibition of binding to membranes of CHO cell line stably transfected with human vasopressin V1a and V2 receptor subtypes, resp. The compds. I are also useful in enhancing fertility rates, enhancing survival rates and synchronizing estrus in farm animals, and may be useful in the prevention and treatment of dysfunctions of the oxytocin system in the central nervous system including obsessive compulsive disorder (OCD) and neuropsychiatric disorders.

MSTR 1

$$G5 = 710$$

$$G6 = N$$

10/572,914

G7 = CONH2 / alkylamino <containing 1-6 C> / SO2NH2

Patent location: claim 1

Note: and pharmaceutically acceptable salts, or prodrugs

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 79 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 137:190369 MARPAT

TITLE: Hair dyes containing cationic quinolinium direct dyes

PATENT ASSIGNEE(S): Wella A.-G., Germany

SOURCE: Ger. Gebrauchsmusterschrift, 25 pp.

CODEN: GGXXFR

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO. I	DATE
DE 20204129	U1	20020829	DE 2002-20204129 2	20020315
PRIORITY APPLN. INFO	١.:		DE 2002-20204129 2	20020315

AB The invention concerns hair dye compns. that contain cationic direct dyes from the group of quinolinium salts. The compns. further contain other direct dyes, e.g. azo dyes, quinone dyes, and triphenylmethanes.

Oxidative dyes, oxidation agent, synthetic polymers or modified natural polymers can be included. Thus 4-[(4-aminophenyl)amino]-1-ethylquinolinium-tetrafluoroborate was synthesized and used at an amount of 0.01 g in a dye that also included 10.00 g ethanol and 10.00 g water. The dye mixture was diluted with 10% citric acid or 10% ammonia solution for testing

the color effects.

MSTR 1

G2 = 17 / SO2NH2

G6 = 76



G7 = heteroaryl

Patent location: claim 1

Note: additional ring formation also claimed

L5 ANSWER 80 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 137:135116 MARPAT

TITLE: Diphenyl ether derivatives, their preparation, and

their uses as heparanase inhibitors

INVENTOR(S): Ayal-Hershkovitz, Maty; Miron, Daphna; Koller, Avi;

Ilan, Neta; Levy, Ofra

PATENT ASSIGNEE(S): Insight Strategy and Marketing Ltd., Israel

SOURCE: PCT Int. Appl., 77 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	CENT 1	. O <i>l</i> .		KI	ND	DATE			APPLICATION NO.					DATE			
		2002		_			2002			M.	0 20	02-I	L82		2002	0129		
	WO	2002	0603	75	Α.	3	2003	1009										
		W:	ΑE,	AG,	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KΖ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
			PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TN,	TR,	TT,	TZ,
			UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW	·	·		·	·	·	·
		RW:	GH,	GM,	KE,	LS,	MW,	MΖ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
			KG,	KΖ,	MD,	RU,	ТJ,	TM,	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FΙ,	FR,	GB,
			GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,
			GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	ΤG							
	ΑU	2002	2300	57	A	1	2002	0812		A	U 20	02-2	3005	7	2002	0129		
PRIO:	RIT:	APP	LN.	INFO	.:					U	S 20	01-2	6430	5P	2001	0129		
										M	0 20	02-1	L82		2002	0129		
\sim T																		

AB The invention provides di-Ph ether compds. as heparanase inhibitors suitable for treatment of diseases and disorders caused by or associated with heparanase catalytic activity, e.g. cancer, inflammatory disorders, and autoimmune diseases. Preparation and biol. activity of e.g. I are described.

MSTR 1

G2 = 31

G8—G9

G8 = C(0) / SO2G14 = 57

G8---G9

G15 = 67

6⁹ 678—G9

G17 = 104 / 116

10/572,914

G18 NH G18 G18 G18 G18 G18 G18

G18 = 127

127—G19

G19 = NH2 (opt. substd.) / heterocycle <containing 5-7 atoms, 1-4 heteroatoms, 1 or more N, zero or more O, zero or more S (no other heteroatoms), attached through 1 or more N, non-aromatic, saturated, 5- to 7-membered monocyclic ring> (opt. substd.)

Patent location: claim 1

Note: and pharmaceutically acceptable salts

L5 ANSWER 81 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 136:85762 MARPAT

TITLE: New aryl-, quinolyl-, and other heterocyclyl-

containing amino alcohol derivatives useful as β 3

adrenergic receptor agonists

INVENTOR(S): Kayakiri, Hiroshi; Sakurai, Minoru; Washizuka,

Kenichi; Hamashima, Hitoshi; Tomishima, Yasuyo; Fujii,

Naoaki; Taniguchi, Kiyoshi

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co., Ltd., Japan

SOURCE: PCT Int. Appl., 121 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

P.	ATENT	NO.		KII	ND	DATE APPLICATION NO.			Э.	DATE							
_																	
M	0 2002	0006	22	A.	2	2002	0020103 WO 2001-JP5425 20					2001	0625				
M	0 2002	0006	22	A.	3	2002	0829										
	W:	ΑE,	AG,	AL,	AM,	ΑT,	ΑU,	ΑZ,	ΒA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KR,	KΖ,	LC,	LK,	LR,	LS,
		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,
		RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,
		VN,	YU,	ZA,	ZW												
	RW:	GH,	GM,	ΚE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,
		DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG		
PRIORI'	TY APP	LN.	INFO	.:					Αl	U 20	8-00	413		2000	0627		
GI																	

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The invention relates to compds. I [wherein: X1 = bond or OCH2; X2 = (CH2)1-2; X3 = bond, O, or NH; R1 = (un)substituted Ph, indolv1, or carbazolyl [substituents = 1 or 2 of OH, halo, NO2, amino, formyl, (lower)alkylsulfonylamino, aryl(lower)alkoxy, and hydroxy(lower)alkyl]; R2 = H or aryl(lower)alkyl; R3 = H or hydroxy(lower)alkyl; R4 = (un) substituted aryl, 4-quinolyl, phthalazinyl, quinazolinyl, cinnolinyl, or naphthyridinyl; with provisos], or their pharmaceutically acceptable salts. The compds. are $\beta 3$ adrenergic receptor agonists, and therefore have gut sympathomimetic, antiulcer, anti-pancreatitis, lipolytic, and smooth muscle relaxant activities. In particular, I and salts are useful for the prophylactic and/or the therapeutic treatment of pollakiuria or urinary incontinence. Sixty precursor prepns. and 63 invention examples, including well over 200 invention compds., are provided. For example, the structure of claimed compound II is typical. Another invention compound, phthalazine derivative III, was prepared from 4-((2S)-2-amino-3-hydroxypropyl)phenol HCl, benzaldehyde, (2S)-3-phenoxy-1,2-epoxypropane, and 1-chlorophthalazine, in 4 steps. III at 0.32 mg/kg (intraduodenal) in beagle dogs gave 35.9% inhibition of carbachol-induced increase in intravesical pressure.

MSTR 1

$$G3 = 497$$

G8 = phenylene

G13 = NH G15 = NH2 G18 = 502

G25 = 654 / 679

10/572,914

Patent location: claim 1

Note: substitution is restricted

Note: and salts

Note: also incorporates claim 5

L5 ANSWER 82 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 136:48475 MARPAT

TITLE: Cationic rhodacyanine dye derivatives as inhibitors

for interaction mot-2 protein and the p53 protein

INVENTOR(S): Wadhwa, Renu; Sugihara, Takashi; Yoshida, Akiko;

Shishido, Tadao

PATENT ASSIGNEE(S): Chugai Bunshi Igaku Kenkyusho K. K., Japan; Fuji Photo

Film Co., Ltd.

SOURCE: Jpn. Kokai Tokkyo Koho, 12 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2001354564 PRIORITY APPLN. INFO.	A :	20011225	JP 2000-184540 JP 2000-184540	20000614 20000614

AB Cationic rhodacyanine dye derivs. (I and II; X1, X2 = S, -CH=CH-; R1, R2,

Ι

10/572,914

R3, R4 = Me, Et; Z1 = -X2-C=(CH-CH)n=N+(R3)- forming ring with thiazole, benzothiazole, thiazolin, 2-pyridine, 2-quinoline, 4-quinoline; q = anion; N = 0, 1) are claimed as inhibitors for interaction mot-2 protein and the p53 protein and are useful for studying cell cycle, cell proliferation, carcinogenesis, and treatment of p53 protein-related diseases, including cancer.

MSTR 1

$$\begin{array}{c} \text{G1} & \begin{array}{c} \text{CH} & \text{G3} & \text{G2} \\ 15 & 35 \end{array} \\ \text{G2} & \begin{array}{c} \text{G4} \end{array}$$

 $G3 = 48-15 \ 47-35$

G5 = acylamino / CONH2 / SO2NH2 Patent location: claim 1

L5 ANSWER 83 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 135:272892 MARPAT

TITLE: Preparation of quinoline derivatives as nuclear

peroxisome proliferator-activated receptors

antagonists

INVENTOR(S): Kadota, Hidetoshi; Fukazawa, Nobuyuki; Nagase,

Hiroshi; Maruyama, Kyoko; Nakao, Toshifumi; Asada, Noriaki; Hachimaki, Toshiyuki; Kibayashi, Kenji; Uta,

Hideyuki; Morikawa, Maki

PATENT ASSIGNEE(S): Mitsui Chemicals Inc., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 39 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2001261654 WO 2001070698	A A1	20010926 20010927	JP 2000-79146 WO 2001-JP2168	20000321 20010319

W: CN, KR, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE, TR

EP 1266888 A1 20021218 EP 2001-914178 20010319

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR

US 20030212100 A1 20031113

US 2002-239310 20020920 JP 2000-79146 20000321

PRIORITY APPLN. INFO.:

WO 2001-JP2168 20010319

GΙ

$$R^2$$
 Y^2 X X^1 Y^2 Y^2 Y^2 Y^2 Y^2 Y^2

AB Title compds. [I; R = (CH3)2CHO, H, CH3O; Y2 = CH, CCH3, N; R2 = H, CH3O, CH3CH2; R3 = H, CH3O, CH3; X = 4-CH2OC6H4CH2CH(OCH2CH3)COOH, 4-CH2OC6H4CH2CH(OCH2CH3)COOCH2CH3, H, 4-CH2OC6H4CH2CH(OC6H5)COOCH3, 4-CH2OC6H4CH2CH(OC6H5)COOH, 4-CH2OC6H4CH2CH(SC6H5)COOCH2CH3, 4-CH2OC6H4CH2CH(OCH2CH3)COOH, 3-CH2OC6H4CH2CH(OCH2CH3)COOCH2CH3, 3-CH2OC6H4CH2CH(OCH2CH3)COOH; X1 = H, 4-CH2OC6H4CH2CH(OCH2CH3)COOCH2CH3, C6H5, 4-CH2OC6H4CH2CH(OCH2CH3)COOH; Y = N, CH; Y1 = CH, N, CC1, CF, CF, etc.] are prepared as PPAR (peroxisome proliferator-activated receptors) antagonists. Title compds. I offer the prevention or treatment of various diseases where PPAR-α and PPAR-γ play roles as the causes. Thus, the title compound II was prepared and biol. tested for PPARα and PPARγ antagonist activities.

ΙI

MSTR 1

G1 = quinolinyl (opt. substd. by 1 or more G2) G2 = 11 / 14

G3 = cycloalkyl <containing 3-4 C> Patent location: claim 1

Note: or pharmacologically acceptable salts

L5 ANSWER 84 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 135:213459 MARPAT

TITLE: Photoelectric converters, photoelectrochemical cells,

and metal complex pigments

INVENTOR(S): Takizawa, Hiroo

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 51 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

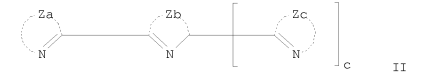
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2001237000	A	20010831	JP 2000-44897	20000222
PRIORITY APPLN. INFO.	:		JP 2000-44897	20000222
O.T.				

GΙ

$$(R_{1})_{a1}$$
 $(R_{1})_{a1}$
 $(R_{1})_{a1}$
 $(R_{1})_{a1}$
 $(R_{1})_{a1}$
 $(R_{1})_{a1}$
 $(R_{2})_{a2}$
 $(R_{1})_{a1}$
 $(R_{2})_{a2}$
 $(R_{2})_{a2}$



AB The photoelec. converters contain semiconductor particles sensitized by a metal complex pigment Lm1Xm2M1L'M2L"m3X'm4.CI, where L' = I, L = single bond, O, S, alkenyl group, alkenylene group, arylene group, or hetero arylene group; R1 = carboxyl sulfonyl, hydroxyl, hydroxamic acid, phosphoryl, or phosphonyl group: R2 = substituents; a1 and a2 = 0-4

integers, R1 can be same or different when al ≥ 2 , and R2 can be same or different or forming a ring when a2 ≥ 2 ; n = 0-2 integer; L and L" = di- or tri-dentate ligand II with Za, Zb, and Zc = nonmetal atoms forming 5- or 6-membered rings, c = 0 or 1; X and X' = mono-or bi-dentate ligand selected from acyloxy, acylthio, thioacyloxy, thioacylthio, acylaminoxy, thiocarbamate, dithiocarbamate, thiocarbonate, dithiocarbonate, trithiocarbonate, acyl, thiocyanate, isothiocyanate, cyanate, isocyanate, cyano, alkylthio, arylthio, alkoxy, aryloxy groups, halogen, carbonyl, dialkyl ketone, 1,3-diketone, carbamide, thiocarbamide, and thiourea; m1 and m3 = 0-2 integers, m2 and m4 = 0-4 integers, X1 and X2 can be same or different or form rings among X1's and/or among X2's when m2 and m4 ≥ 2 ; and CI = charge balancing counter ions. Photoelectrochem. cells use the photoelec. converters.

MSTR 1

G5 G3 G21

G3 = 483

G8 = 477 / 574 / 572

 $\begin{array}{lll} {\rm G10} & = & {\rm NH2} \\ {\rm G13} & = & {\rm NH} \\ {\rm G14} & = & {\rm acyl} \\ {\rm G15} & = & {\rm NH2} \\ \end{array}$

Patent location: claim 1

Note: additional ligands also claimed

Note: as complexes

Note: substitution is restricted

L5 ANSWER 85 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 135:197978 MARPAT

TITLE: Photoelectrochemical cells

INVENTOR(S):
Takizawa, Hiroo

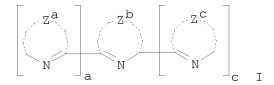
PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 25 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2001229983	A	20010824	JP 2000-37290	20000215
PRIORITY APPLN. INFO.	:		JP 2000-37290	20000215
GI				



AB The cells use semiconductor particles sensitized by metal complex pigments M(NR1R2R3)mLm'.CI, where M = metal atom, R1-3 = H, alkyl, alkenyl or aryl groups, L = 1-3 dentate ligand I (Z1, Z2, Z3 = non-metal atoms forming 5- or 6-membered rings, p and q = 0 or 1), m = 1-5, (NR1R2R3) can be different from each other or joined together when m \geq 2, m' = 1 or 2, L can be different from each other when m' =2, and CI = counter ion for elec. balance of the pigment.

MSTR 1

$$G3 = 483$$

$$G8 = 477 / 574 / 572$$

 $\begin{array}{lll} {\rm G10} & = & {\rm NH2} \\ {\rm G13} & = & {\rm NH} \\ {\rm G14} & = & {\rm acyl} \\ {\rm G15} & = & {\rm NH2} \\ \end{array}$

Patent location: claim 1

Note: additional ligands also claimed

Note: as complexes with G5

Note: substitution is restricted

L5 ANSWER 86 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 135:76882 MARPAT

TITLE: Preparation of heterocyclic compounds having

sulfonamide groups as inhibitors of angiogenesis INVENTOR(S): Haneda, Toru; Tsuruoka, Akihiko; Kamata, Junichi;

Okabe, Tadashi; Takahashi, Keiko; Nara, Kazumasa; Hamaoka, Shinichi; Ueda, Norihiro; Wakabayashi, Toshiaki; Funahashi, Yasuhiro; Semba, Taro; Hata,

Naoko; Yamamoto, Yuji; Ozawa, Yoichi; Tsukahara, Naoko; Owa, Takashi

PATENT ASSIGNEE(S): Eisai Co., Ltd., Japan

SOURCE: PCT Int. Appl., 94 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	ENT	NO.		KII	ND	DATE			AI	PPI	LICA	ATI(ои ис	Э.	DATE			
															20001	1227		
						JP,												
	RW:					DE,	DK,	ES,	FΙ,	FF	₹, (GΒ,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,
		PT,	SE,	TR														
CA	2395	772		A.	1	2001	0705		CI	A 2	2000	0-23	3957	72	20003	1227		
ΑU	2001	0222	83	A		2001	0709		JA	J 2	2001	1 - 22	2283		20001	1227		
ΑU	7769	33		B	2	2004	0923											
									EF	2	2000)-98	35953	3	20003	1227		
EP	1243	583		B	1	2005	0928											
									GB,	GF	₹, :	ΙΤ,	LI,	LU,	NL,	SE,	MC,	PT,
			FΙ,															
HU	2002	0039	73	A.	2	2003	0328		JН	J 2	2002	2-39	973		20003	1227		
HU	2002	0039	73	A.	3	2004	0728											
NΖ	5193	80		А		2004	1029		NZ	3 2	2000	0-51	19380	0	20003	1227		
RU	2239	631		C	2	2004	1110		RU	J 2	2002	2-12	2051	5	20001	1227		
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ES	2246	922		T	3	2006	0301		ES	3 2	2000	0-98	35953	3	20001 20001 20001 20001	1227		
US	2003	0144	507	A:	1	2003	0731		US	3 2	2002	2-14	49253	3	20020	0610		
NO	2002	0030	97	A		2002	0828		NO) 2	2002	2-30	97		20020	1626		
NO	3242	68		B	1	2007	0917			_								
MX	2002	PA06	474	Α.	_	2002	1129		MΣ	7 2	2002	2-P	4647	4	20020	0627		
	APP														1999			
				• •											20001			
									000		_ 000	0 01	. , , , ,	0	2000.	1		

OTHER SOURCE(S): CASREACT 135:76882

GI

AB Heterocyclic compds. having sulfonamide or sulfonylurea groups, specifically heterocyclic compds. of general formula (I), pharmacol. acceptable salts of the same, or hydrates of both [wherein A is hydrogen, halogeno, optionally halogenated C1-4 alkyl, hydroxy, cyano, (CO)kNR2R3, or optionally substituted C2-4 alkenyl or alkynyl (wherein R2 and R3 are each independently hydrogen or optionally halogenated C1-4 alkyl; k is 0 or 1); B is optionally substituted aryl, monocyclic heteroaryl, or Q1 (wherein the ring Q is an optionally substituted aromatic ring containing 1 or 2

N atoms; the ring M is optionally substituted and unsatd. C5-12 monocyclic or polycyclic ring sharing a double bond with the ring Q and optionally containing 1-4 heteroatom selected from N, O, and S; the ring Q and M may share a N atom); K is a single bond or (CR4R5)m (wherein R4 and R5 are each independently hydrogen or C1-4 alkyl; m is 1 or 2); T, W, X and Y are each independently =C(D)- (wherein D is hydrogen, halogeno, hydroxy, C1-4 alkyl, halo-C1-4 alkyl, or the like) or nitrogen; U and V are each independently =C(D)-, nitrogen, oxygen, or CO; Z is a single bond or -CONH-; and R1 is hydrogen or C1-4 alkyl] are prepared These compds. includes N-quinolinylpyridinesulfonamides, N-quinolinylbenzenesulfonamides , N-quinolinylquinolinesulfonamides, N-quinolinylindolinesulfonamides, N-quinolinylisoquinolinesulfonamides, N-quinolinylbenzofuransulfonamides, N-quinolinyltetrahydronaphthalanesulfonamides, Nquinolinylbenzoxathiansulfonamide, N-quinolinylbenzothiopyransulfonamide, N-isoquinolinylpyridinesulfonamides, N-isoquinolinylbenzenesulfonamides, N-naphthyridinylpyridinesulfonamides, N-naphthyridinylbenzenesulfonamides, N-quinolinylpyridazinesulfonamides, etc. They are useful as therapeutics based on angiogenesis inhibition such as antitumor agents, cancer metastasis inhibitors, and therapeutics for diabetic retinopathy, rheumatic arthritis, and hemangioma. Thus, 5-indansulfonyl chloride was added to a solution of 3-amino-8-bromoquinoline in pyridine and stirred at room temperature for 30 min to give N-(8-bromoquinolin-3-y1)-5-indansulfonamide (II). II and N-(8-bromoquinolin-3-yl)-6-methoxypyridazine-3-sulfonamide in vitro showed IC50 of 0.04 and 0.53 μ g/mL, resp., against angiogenesis in rat aorta.

MSTR 1

G13 = bondG14 = N / 38

с——G15 38

G15 = alkylamino < containing 1-4 C>

(opt. substd. by 1 or more G2) / CONH2

 $G16 = 42-5 \ 43-8$



Patent location: claim 1

Note: substitution is restricted

Note: additional ring formation also claimed

REFERENCE COUNT: 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 87 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 134:252257 MARPAT

TITLE: Preparation of 2-(indolin-3-yl)quinoline derivatives

and compositions in use as antimicrobial agents

INVENTOR(S): Cuny, Gregory D.; Hauske, James R.; Heefner, Donald

L.; Hoemann, Michael Z.; Kumaravel, Gnanasambandam;

Melikian-Badalian, Anita; Rossi, Richard F.

PATENT ASSIGNEE(S): Sepracor, Inc., USA

SOURCE: U.S., 112 pp., Cont.-in-part of U.S. Ser. No. 878,781,

abandoned.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 7

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6207679	В1	20010327	US 1998-45051	19980319
CA 2293418	A1	19981223	CA 1998-2293418	19980618
WO 9857931	A2	19981223	WO 1998-US12762	19980618
WO 9857931	A3	19990429		

W: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK,

GΙ

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EE, ES, FI, GB, GE, GH, BM, GW, HU, ID, IL, IS, JP, KE, KG, KP,
             KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MN, MW, MX, NO, NZ,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG,
             US, UZ, VN, YU, ZW
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,
             FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,
                     GN, ML, MR, NE, SN, TD, TG
     EP 991623
                       Α2
                           20000412
                                            EP 1998-930396
                                                              19980618
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
     US 6172084
                        В1
                             20010109
                                            US 1998-99640
                                                              19980618
     HU 2000003364
                        Α2
                             20010628
                                            HU 2000-3364
                                                              19980618
     HU 2000003364
                       АЗ
                             20020328
     JP 2002505689
                        Τ
                             20020219
                                            JP 1999-504835
                                                              19980618
     AU 757059
                       В2
                                            AU 1998-79797
                             20030130
                                                              19980618
     US 6103905
                                            US 1998-213385
                                                              19981211
                        Α
                             20000815
     NO 9906269
                                            NO 1999-6269
                             20000216
                                                              19991217
                        Α
     US 6376670
                             20020423
                                            US 2000-658690
                                                              20000908
                        В1
PRIORITY APPLN. INFO.:
                                            US 1997-878781
                                                              19970619
                                            US 1998-45051
                                                              19980319
                                            US 1998-99640
                                                              19980618
                                            WO 1998-US12762
                                                              19980618
                                            US 1998-213385
                                                              19981211
                                            US 2000-639622
                                                              20000815
```

$$R^4$$
 A
 R^3
 R^3
 R^3
 R^3
 R^3
 R^3
 R^3
 R^3

F3C NH2

AB Title compds. I [wherein; R, R1, R2 and R3 are H, halo, alk(en)(yn)yl, OH,

Ι

alkoxy, amino, nitro, SH, imine, amide, CO, -(CH2)0-8-R80, etc.; R4 is the same as R-R3 but not H; R5 is the same as R4 except that at least 1(-8) CH2 precede R80; A is (un)substituted with any number of R4 up to the number limited by stability and rules of valence; B is substituted with at least one instance of R5 up to the number limited by stability and rules of valence; R80 is (substituted) aryl, cycloalk(en)yl, heterocyclyl or polycyclyl.] and related quinoline derivs. are prepared as antimicrobial agents. For instance, synthesis of II is accomplished by alkylation of 4-hydroxymethyl-6-trifluoromethyl-2-(N-t-butoxycarbonylindol-3-yl)quinoline with (4-t-butoxycarbonylaminomethyl)benzyl iodide followed by deprotection. There are 282 examples of I provided. The min. inhibitory concentration (MIC) of I against at least one Gram-pos. bacterium is 0.1-10 μ g/mL. Certain compds. of formula I have a therapeutic index in primates of at least 10 for the inhibition of infection by at least one Gram-pos. bacterium.

MSTR 1

G1 = o-C6H4 (opt. substd. by G2) G2 = 22

c(0)-G9

G9 = NH2 (opt. substd.) / heterocycle <containing 4-8
 atoms, 1 or more N, attached through 1 or more N>
 (opt. substd.)

G21 = N G23 = 98

g(O)—G9

G28 = NHC(NH)NH2 (opt. substd.)
Patent location: claim 1

Note: also incorporates later claims and broader

disclosure

REFERENCE COUNT: 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 88 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 134:86170 MARPAT

TITLE: Quinoline-indole antimicrobial agents

INVENTOR(S): Cuny, Gregory D.; Hauske, James R.; Heefner, Donald L.; Hoemann, Michael Z.; Kumaravel, Gnanasambandam;

Melikian-badalian, Anita; Rossi, Richard F.

PATENT ASSIGNEE(S): Sepracor, Inc., USA

SOURCE: U.S., 151 pp., Cont.-in-part of U.S. Ser. No. 45,051.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 7

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6172084 US 6207679 US 6103905 US 6376670 PRIORITY APPLN.	B1 B1 A B1 INFO.:	20010109 20010327 20000815 20020423	US 1998-99640 US 1998-45051 US 1998-213385 US 2000-658690 US 1997-878781 US 1998-45051 US 1998-99640 US 1998-213385 US 2000-639622	19980618 19980319 19981211 20000908 19970619 19980319 19980618 19981211 20000815

Ι

GΙ

$$R^4$$
 R^3
 R^6
 R^7
 R^7

$$\begin{array}{c|c} & \text{H}_2\text{C}-\text{O}-\text{CH}_2 \\ \hline \\ \text{F}_3\text{C} \\ \hline \\ \text{N} \\ \\ \text{H} \end{array} \\ \text{Br}$$

AB Indolylquinolines I [X = N; Y = NR; R-R3 = independently H, halogen, alkyl, alkenyl, alkynyl, OH, alkoxy, silyloxy, NH2, NO2, SH, alkylthio, imino, amido, phosphoryl, phosphonate, phosphine, CO, CONH2, anhydride, silyl, alkylsulfonyl, arylsulfonyl, alkylseleno, aldehyde, ester, heteroalkyl, CN, guanidine, amidine, acetal, ketal, amine oxide, (hetero)aryl, azide, aziridine, carbamate, epoxide, C(:NH)OH, imide, oxime, SO2NH2, CSNH2, thiocarbamate, urea, thiourea, or (CH2)mR80; R4R5, R6R7 = atoms required to complete an (un)substituted fused benzo ring

II

system; R80 = (un)substituted aryl, cycloalkyl, cycloalkenyl, heterocycle, or polycycle; m = 0-8] were prepared by conventional or combinatorial synthetic methods for use as bactericides. Thus, 4-H2NCH2C6H4C02H was esterified, N-tert-butoxycarbonylated, reduced, and treated with iodine to give 4-BocNHCH2C6H4CH2I, which was coupled with the indolylquinolinemethanol fragment and deblocked to give the product II. II had MIC's <7 $\mu g/mL$ against methicillin-resistant Staphylococcus aureus, vancomycin-resistant Enterobacter sp., and Streptococcus pneumoniae.

MSTR 1

G1 = o-C6H4 (opt. substd. by G2)

G2 = 22

2C(O)—G9

G9 = NH2 (opt. substd.) / heterocycle <containing 4-8 atoms, 1 or more N, attached through 1 or more N>

(opt. substd.)

G21 = N G23 = 98

98 (O)—G9

G28 = NHC(NH)NH2 (opt. substd.)
Patent location: claim 1

Note: also incorporates later claims and broader

disclosure

REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 89 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 133:245161 MARPAT

TITLE: Rewritable optical recording materials containing azo

chelates

INVENTOR(S): Ueno, Yasunobu; Sato, Tsutomu; Tomura, Tatsuya; Azuma,

Yasuhiro

PATENT ASSIGNEE(S): Ricoh Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 15 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2000263942	A	20000926	JP 1999-75978	19990319
JP 3682759	В2	20050810		
PRIORITY APPLN. INFO.	:		JP 1999-75978	19990319
GI				

AB The recording layer of the materials contain azo chelates comprising of azo compound I (R1-2 = H, (un)substituted alkyl, aryl; R1 and R2 may form a ring; R3-11 = H, halogen, nitro, cyano, OH, carboxyl, amino, carbamoyl, (un)substituted alkyl, aryl, heterocycle, etc; 2 of the neighboring R3-11 may form rings; X = OH, alkyloxy, aryloxy, carboxy, amino, sulfo, etc.) and a metal. The materials are resistant to light and are storage stable.

Ι

MSTR 1

G3 = CONH2 / alkylcarbonylamino (opt. substd.)

Patent location: claim 1

Note: as metal chelates

Note: additional ring formation also claimed

L5 ANSWER 90 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 132:265101 MARPAT

TITLE: Preparation of 3-cyanoquinolines as protein tyrosine

kinase inhibitors

INVENTOR(S): Wissner, Allan; Tsou, Hwei-Ru; Berger, Dan Maarten;

Floyd, Middleton Brawner, Jr.; Hamann, Philip Ross;

Zhang, Nan; Salvati, Mark Ernest; Frost, Philip

PATENT ASSIGNEE(S): American Cyanamid Company, USA

SOURCE: PCT Int. Appl., 195 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	TENT :							APPLICATION NO.			DATE						
WO	2000 W:	AE, CZ, IN, MG, SL, GH, DK,	AL, DE, IS, MK, TJ, GM, ES,	AM, DK, JP, MN, TM, KE, FI,	AT, DM, KE, MW, TR, LS,	2000 AU, EE, KG, MX, TT, MW, GB,	AZ, ES, KP, NO, TZ, SD, GR,	BA, FI, KR, NZ, UA, SL, IE,	BB, GB, KZ, PL, UG, SZ, IT,	D 19 BG, GD, LC, PT, UZ, TZ, LU,	99-U BR, GE, LK, RO, VN, UG,	S220 BY, GH, LR, RU, YU, ZW, NL,	CA, CA, GM, LS, SD, ZA, AT,	CH, HR, LT, SE,	CN, HU, LU, SG,	ID, LV, SI,	IL, MD, SK,
AU AU EP	2344 9961 7636 1117 1117	169 593 69 659		A A B	1 2 1	2000 2000 2003	0406 0417 0731 0725		C. Al	A 19 U 19	99-2. 99-6	3441 1593	69	1999 1999 1999	0922		
HU JP NZ AT PT ES SK TW NO NO MX IN ZA HK	2001 2002 5105 2555 1117 2211 2848 2334 2001 3245 2001 2001 1035 2007	IE, 0035 0035 5253 51 75 659 175 46 37 0015 63 PA03 KN00 0027 188 KN02	SI, 20 20 69 75 230 370 29	LT, A. A. A. A. A. A.	LV, 2 3 3	FI, 2002 2003 2002 2003 2004 2004 2005 2005 2001 2007 2001 2006 2002	RO 0228 0128 0128 0328 1215 0430 0701 1201 0601 0528 1119 1011 0303 0703 0402		HI N' A' P' E: SI TV NO	U 20 P 20 Z 19 T 19 T 19 S 19 K 20 W 19 O 20 X 20 N 20 K 20 N 20 K 20 N 20 N 20	01-3 00-5 99-5 99-9 99-9 99-9	7222 1055 4841 4841 13 8116 575 A323 N370 729 0582 N234	1 1 0 0 0 0 0 6 3 0	NL, 1999 1999 1999 1999 1999 1999 2001 2001	0922 0922 0922 0922 0922 0922 0922 0923 0328 0328 0329 0403 0817 0625	MC,	PT,
GI			11VI O	• •					M	0 19		S220	54	1999 2001	0922		

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AB X(CH2)nZZ1CN [I; X = (un)substituted bicyclic (hetero)aryl or LTA; A = (un)substituted phenylene, -pyridinediyl, -pyrimidinediyl; T = O, S, (alkyl)imino(alkylene), oxyalkylene, etc.; Z = O, S, (alkyl or alkanoyl)imino; Z1 = 2-unsubstituted-5,6,7,8-(un)substituted quinoline-4,3-diyl; n = O or 1] were prepared Thus, Me 2-amino-4,5-diethoxybenzoate was N-condensed with HCNMe2/POC13 and the product cyclocondensed with MeCN to give, after POC13 treatment, 4-chloro-6,7-diethoxyquinoline-3-carbonitrile which was aminated by 6-aminoindoline to give title compd II. Data for biol. activity of I were given.

MSTR 1

$$\begin{array}{c|c} & G31 \\ 60 & |57 \\ & G10 - G2 - G1 \\ & & G32 \\ & & & G31 \\ & & & & & \\ & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & & \\ & & & \\ & & \\ & &$$

G1 = 51

$$G2 = NH$$
 $G8 = 22-12 21-52$

 $G10 = 77-12 \ 81-57 \ 80-60 \ 79-59 \ 78-58$

G32 = alkylaminocarbonyl

G41 = N

Derivative: or pharmaceutically acceptable salts

Patent location: claim 1

Note: substitution is restricted also incorporates claim 16

Note: additional ring formation also claimed

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 91 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 132:265100 MARPAT

TITLE: Preparation of substituted 3-cyanoquinolines as

protein tyrosine kinases inhibitors

INVENTOR(S): Wissner, Allan; Tsou, Hwei-Ru; Berger, Dan Maarten;

Floyd, Middleton Brawner, Jr.; Hamann, Philip Ross;

Zhang, Nan; Frost, Philip

PATENT ASSIGNEE(S): American Cyanamid Company, USA

SOURCE: PCT Int. Appl., 164 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE						
WO 2000018740	A1 20000406	000406 WO 1999-US22056 19990922							
	AM, AT, AU, AZ, BA,								
	DK, DM, EE, ES, FI,								
· · · · ·	JP, KE, KG, KP, KR,		· · · · · · · · · · · · · · · · · · ·						
· · ·	MN, MW, MX, NO, NZ,								
· · · · ·	TM, TR, TT, TZ, UA,								
· · ·	KE, LS, MW, SD, SL,								
· ·	FI, FR, GB, GR, IE,		· · · · · ·						
	CM, GA, GN, GW, ML,		51, 51, 50, 61,						
	A1 20000406		19990922						
	A 20000417								
	A 20010626								
	A1 20010025 A1 20010725								
· · ·	CH, DE, DK, ES, FR,	GB, GK, 11, L1, LU,	NL, SE, MC, PI,						
	LT, LV, FI, RO	0001 0600	1000000						
	A2 20020228	HU 2001-3633	19990922						
	A3 20030128								
	T 20020813								
NZ 510580	A 20030328	NZ 1999-510580	19990922						
	A1 20080730								
R: AT, BE,	CH, CY, DE, DK, ES,	FI, FR, GB, GR, IE,	IT, LI, LU, MC,						

NL, PT,	SE, AL	L, LT, LV, RO,	SI		
ZA 2001002501	A	20020105	ZA	2001-2501	20010327
NO 2001001574	A	20010528	NO	2001-1574	20010328
MX 2001PA03227	A	20011011	MX	2001-PA3227	20010328
AU 2004200300	A1	20040219	AU	2004-200300	20040128
AU 2007201934	A1	20070524	AU	2007-201934	20070501
PRIORITY APPLN. INFO.	:		US	1998-162289	19980929
			AU	1999-61594	19990922
			EP	1999-948411	19990922
			WO	1999-US22056	19990922
			AU	2004-200300	20040128
O.T.					

GΙ

$$R^1$$
 Z (CH₂) nX CN G^2 N

Ι

AB The title compds. I [X = cycloalkyl, pyridinyl, pyrimidinyl, etc.; Z = NH, O, S, NR; G1, G2, R1, R4 = H, halo, alkyl, alkynyl, etc.; n = 0,1], protein tyrosine kinase inhibitors, were prepared E.g., 4-(2-methoxyethoxy) but-2-ynoic acid [4-(3-bromophenylamino)-3-cyanoquinolin-6-yl]amide was prepared I are useful as antineoplastic agents.

MSTR 1

$$\begin{array}{c} & G31 \\ 60 & 57 \\ 60 & G10 - G2 - G1 \\ G32 & 56 & 12 & 13 \\ & & & & & \\ 58 & & & & & \\ & & & & & \\ 58 & & & & & \\ \end{array}$$

$$G1 = 22$$

G2 = NHG10 = 88-12 92-57 90-60 91-59 89-58

```
C(0)-NH<sub>2</sub>
```

= 99 / 112 / 115 G11

9G17-G14 G12-G17-G18 G17-G18

= alkylene <containing 1 or more C> (opt. substd.) G17

= alkylaminocarbonyl / 246 / 248 / 251 G32

G12—G17—G18 G17—G18

G41 = N

Derivative: or pharmaceutically acceptable salts

Patent location: claim 1

Note: substitution is restricted Note: also incorporates claim 10

Note: additional ring formation also claimed

REFERENCE COUNT: THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS 4 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 92 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 132:175808 MARPAT

TITLE: Hepatitis C inhibitor peptides

INVENTOR(S): Llinas-Brunet, Montse; Bailey, Murray D.; Cameron, Dale; Ghiro, Elise; Goudreau, Nathalie; Poupart, Marc-Andre; Rancourt, Jean; Tsantrizos, Youla S.

PATENT ASSIGNEE(S): Boehringer Ingelheim (Canada) Ltd., Can.

PCT Int. Appl., 113 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			ND	DATE			A	PPLI	CATI	ON No	٥.	DATE			
							_								
WO 2000009	558	А	1	2000	0224		W	0 19	99-C.	A737		1999	0809		
W: AE	, AL,	ΑM,	ΑT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,
DE	, DK,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,
JP	KE,	KG,	KΡ,	KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,
MN	, MW,	MX,	NO,	NΖ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,
TM	TR,	TT,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW					
RW: GH	, GM,	KΕ,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	DE,	DK,
ES	, FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ΒJ,	CF,	CG,
CI	, CM,	GΑ,	GN,	GW,	ML,	MR,	ΝE,	SN,	TD,	ΤG					
US 6767991		В	1	2004	0727		U	S 19	99-3	6867	0	1999	0805		

	2336597 2336597		20000224 20060214		CA	1999-233659	7 1999	0809		
	9952732	A	20000214		AU	1999-52732	1999	0809		
AU	764655	В2	20030828							
BR	9912943	A	20010508		BR	1999-12943	1999	0809		
EP	1105422	A1	20010613		EP	1999-938085	1999	0809		
EP	1105422	B1	20060215							
	R: AT, BE,	CH, DE,	DK, ES,	FR,	GB, C	GR, IT, LI, I	LU, NL,	SE,	MC,	PT,
	IE, SI,	LT, LV,	FI, RO,	CY						
TR	200100438	T2	20010621		TR	2001-438	1999	0809		
HU	2001004548	A2	20020429		HU	2001-4548	1999	0809		
HU	2001004548	A3	20021228							
JP	2002522557	T	20020723		JP	2000-565004	1999	0809		
EE	200100080	A	20020815		EE	2001-80	1999	0809		
NZ	510395	A	20031219		NZ	1999-510395	1999	0809		
TW	577895	В	20040301		TW	1999-881135	37 1999	0809		
	317854		20060315			1999-938085				
	2257066		20060716			1999-938085				
	2001000604	A	20010205			2001-604				
	2001000972	A	20020718			2001-972				
	2001PA01422	A	20000821			2001-PA1422				
	2001MN00128	A	20050304			2001-MN128		-		
	105230	A	20011031		BG	2001-105230	2001	0208		
	64956	B1	20061031							
	2001000101		20020228			2001-101				
	1039947		20050225			2002-101472				
PRIORIT	Y APPLN. INFO	.:				1998-95945P				
						1997-55186P				
						1998-131758				
						1998-219939				
_					WO	1999-CA737	1999	0809		

GΙ

AB The invention provides peptides I (a, b = 0, 1; Y = H, C1-6 alkyl; B = H, acyl derivative, sulfonyl derivative; W = OH, N-substituted amino), or a pharmaceutically acceptable salt or ester thereof, for use in the treatment of hepatitis C virus infection. Preparation of peptides is included.

MSTR 1

G18 = 57

G23-G19

G19 = quinolinyl (opt. substd. by (1-2) G34)

G34 = CONH2 / dialkylamino <each alkyl containing 1-6 C>

Derivative: or pharmaceutically acceptable salts or esters

Patent location: claim 1

Stereochemistry: 32,36,39 - D,L

Stereochemistry: and racemates, diastereoisomers and optical isomers

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 93 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 132:100245 MARPAT

TITLE: Organic electroluminescent device

INVENTOR(S): Takano, Akiko; Himeshima, Yoshio; Tominaga, Takeshi

PATENT ASSIGNEE(S): Toray Industries, Inc., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 2000012223 A 20000114 JP 1998-178373 19980625

PRIORITY APPLN. INFO.: JP 1998-178373 19980625

GI

10/572,914

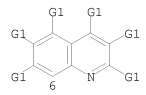
$$R^4$$
 R^3 R^2 R^6 N R^1 OLi I

AB The invention relates to an organic electroluminescent device comprising the 8-hydroxyquinone lithium complex represented by I [R1-6 = H, alkyl, cycloalkyl, etc.].

MSTR 1

HO-G4 Li

G1 = CONH2 / NMe2G4 = 6



Patent location: claim 1

Note: additional substitution also claimed

L5 ANSWER 94 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 131:163194 MARPAT

TITLE: Quinolinol derivative, quinolinol derivative-metal

complex, and organic electroluminescent device

containing it

INVENTOR(S): Ichinosawa, Akiko; Sato, Yoshiharu

PATENT ASSIGNEE(S): Mitsubishi Chemical Industries Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 26 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11204260		19990730	JP 1998-7583	19980119
JP 3772506	R R2	20060510	JP 1990-7303	19900119
PRIORITY APPLN. INFO.		20000310	JP 1998-7583	19980119
GT				

AB The claimed quinolinol derivative and its metal complex have structure I and II, resp. [Ar1-2 = (substituted) aromatic (heterocyclic) group; R1-5 = H, halo, cyano, NH3, NO2, CO2H, OH, (substituted) alkyl, aralkyl, alkenyl, alkynyl, secondary or tertiary amino, amido, acyl, alkoxycarbonyl, alkoxy, alkylsulfonyl, aromatic hydrocarbon group, or aromatic heterocyclic group; R1 and R2, R2 and R3, or R4 and R5 may form ring; M = Be, Zn, Cd, Al, Ga, In, Sc, Y, Mg, Ca, Sr, Co, Cu, Ni, Sm, Eu, Si, Ge, Sn, Tb; n = 2-4]. The electroluminescent device containing the metal complex, preferably in an anode buffer layer formed between an anode and a hole-transporting layer, is also claimed. The electroluminescent device stably emits light in high luminescent efficiency with low driving voltage.

MSTR 1

G2 = CONH2 (opt. substd.) / acylamino

Patent location: claim 1

Note: additional ring formation also claimed

Note: also incorporates claim 2

L5 ANSWER 95 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 130:168399 MARPAT

TITLE: Preparation of ring-bridged bis-quinolines for the

treatment of degenerative diseases of the central

nervous system

INVENTOR(S): Schohe-Loop, Rudolf; Seidel, Peter-Rudolf; Bullock,

William; Feurer, Achim; Terstappen, Georg;

10/572,914

Schuhmacher, Joachim; Vander Staay, Franz-Josef; Schmidt, Bernard; Fanelli, Richard J.; Chisholm, Jane

C.; McCarthy, Richard T.

PATENT ASSIGNEE(S): Bayer A.-G., Germany

SOURCE: U.S., 14 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 5866562 A 19990202 US 1996-738123 19961025

PRIORITY APPLN. INFO: US 1996-738123 19961025

OTHER SOURCE(S): CASREACT 130:168399

GΙ

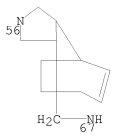
* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The title compds. [I; A, A1, D, D1, E, E1, G, G1, L, L1 = H, cyclopropyl, cyclopentyl, etc.; R1R2 = II-IV (wherein R5, R7 = H, Ph, cyclopentyl, etc.; R6 = H, Me; b = 1-3; R8, R9 = H; or R8 = H, and R9 = R5), etc.] and their salts, useful for the treatment of degenerative diseases such as dementia, were prepared Thus, general procedure for preparing bis-quinolines I was given. E.g., compound V showed Ki of 35 nM/L against 125-apamine binding to bovine cerebral membranes and 73% inhibition of the Rb efflux at 10 μ M.

MSTR 1

$$G1 = 41$$

G3 = 56-7 67-13



Patent location: claim 1

Note: substitution is restricted

REFERENCE COUNT: 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 96 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 129:177224 MARPAT

TITLE: Pickling accelerators, pickling liquid composition

containing them, and pickling method for metal using

the composition

INVENTOR(S): Sasaki, Hiroshi; Okahara, Haruo; Fujiwara, Kazushi

PATENT ASSIGNEE(S): Asahi Chemical Co., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 14 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 10183186	A	19980714	JP 1996-346245	19961225
JP 4028014	В2	20071226		

PRIORITY APPLN. INFO.: JP 1996-346245 19961225

AB A pickling composition comprises at least one compound selected from formic acid,

metal formates, compds. derived by neutralizing formic acid, N-containing heterocyclic compds. (in particular optionally substituted pyridine, quinoline, and isoquinoline), and compds. derived by neutralizing N-containing heterocyclic compds. This pickling method substantially shortens time required for removing surface oxide coatings or contaminants without lowering color tone or increase in corrosion of base metals, does not require equipments for removing poisonous gas, and does not lower quality of base metals such as steel in the recycling step. Thus, 1 g formic acid was added to a solution of 50 g Fe2+ ions and 100 g HCl in 1 L H2O to give a pickling acid liquid The liquid was warmed to 80°, in which a hot rolling steel plate attached with mill scale was immersed. It took 16.2 s to remove mill scale and surface rust vs. 20.5 s without adding the pickling accelerator.

MSTR 2

G1 = NHNH2 / CONH2

Patent location: claim 3

L5 ANSWER 97 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 129:69033 MARPAT

TITLE: Multicomponent system for altering, degrading, or

bleaching lignin, lignin-containing materials, or

similar substances, and method for its use

INVENTOR(S): Freudenreich, Johannes; Stohrer, Juergen; Amann,

Manfred; Mueller, Robert

PATENT ASSIGNEE(S): Consortium fuer Elektrochemische Industrie G.m.b.H.,

Germany

SOURCE: Ger. Offen., 12 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	TENT NO.		KIND	DATE		APPLICATION NO.	DATE			
DE	19651099		A1	19980610		DE 1996-19651099	19961209			
CA	2271937		A1	19980618		CA 1997-2271937	19971205			
WO	9826127		A1	19980618		WO 1997-EP6802	19971205			
	W: AU,	BR,	CA, CN	, JP, KR,	NO,	PL, RU, UA, US				
	RW: AT,	BE,	CH, DE	, DK, ES,	FΙ,	FR, GB, GR, IE, IT	, LU, MC,	NL,	PT,	SE
AU	9855603		A	19980703		AU 1998-55603	19971205			
AU	719140		В2	20000504						
EP	943032		A1	19990922		EP 1997-952038	19971205			
EP	943032		В1	20000913						
	R: AT,	DE,	ES, SE	, PT, FI						
CN	1240008		A	19991229		CN 1997-180387	19971205			
BR	9714387		A	20000516		BR 1997-14387	19971205			
JP	200050584	14	T	20000516		JP 1998-526185	19971205			
			C1	20000820		RU 1999-114460	19971205			
AT	196331			20000915		AT 1997-952038	19971205			
ES	2150797		Т3	20001201		ES 1997-952038	19971205			
PT	943032		T	20001229		PT 1997-952038	19971205			
PRIORITY	Y APPLN. I	NFO.	. :			DE 1996-19651099	19961209			
						WO 1997-EP6802	19971205			

AB The title compns., especially useful in cellulose pulp manufacture, contain oxidants,

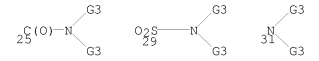
mediators (hydroxylated heterocyclic amines bearing NO or SH groups or their derivs.), and optionally, oxidation catalysts. Adding 20 mL H2O containing

65.3 mg 8-hydroxy-5-nitrosoquinoline (acidified to pH 4.5) and 5 mL H20

containing 15 units of laccase (from Trametes versicolor) to 5 g (dry basis) delignified softwood pulp, kneading for 2 min, and holding in 0 at $45^{\circ}/1-10$ bar for 1-4 h gave pulp with lignin degradation 11.6%.

MSTR 2

G1 = 25 / 29 / 31



G3 = Ph

Derivative: and tautomers, salts, ethers or esters

Patent location: claim 1

Note: additional ring formation also claimed

L5 ANSWER 98 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 129:27902 MARPAT

TITLE: Preparation of bisquinoline compounds for the

treatment of cerebral disorders

INVENTOR(S): Schohe-Loop, Rudolf; Seidel, Peter-rudolf; Bullock,

William; Feurer, Achim; Terstappen, Georg; Schuhmacher, Joachim; Vander Staay, Franz-josef;

Schmidt, Bernard; Fanelli, Richard J.; Chisholm, Jane

C.; McCarthy, Richard T.

PATENT ASSIGNEE(S): Bayer A.-G., Germany

SOURCE: U.S., 18 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 5756517 A 19980526 US 1996-738124 19961025

PRIORITY APPLN. INFO.: US 1996-738124 19961025

AB The title compds. [I; R1, R2 = Me, H; A, A' = H, C1, Me, OMe, etc.; D, D' = H, Me; E, E' = denote hydrogen; G, G' = H; LL' = HN(CH2)2CHEtNH] are prepared I are useful for the treatment of cerebral disorders (no data). Thus, 4-chloro-2-methylquinoline was reacted with H2N(CH2)2CHEtNH2 at 160° for 16 h and then treated with aqueous NaOH to give I (R1 = R2 = Me, A = A' = D = D' = E = E' = G = G' = H, LL' = HN(CH2)2CHEtNH).

MSTR 1

G1 = CONH2G2 = 32-7 34-13

_G3---G5---G3 32---34

G3 = NH

G5 = cyclohexylene

Patent location: claim 2

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 99 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 127:324494 MARPAT

TITLE: Novel polyhalomethane compound and photosensitive

material using it

INVENTOR(S): Okada, Hisashi

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 14 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 09244177	А	19970919	JP 1996-47205	19960305
PRIORITY APPLN. INFO.	:		JP 1996-47205	19960305
GT				

AB The polyhalomethane compound I (R1-7 = H, substituent; ≥ 1 of R2-7 = YCAX1X2; Y = CO, SO, SO2; X1-2 = halo; A = H, electron withdrawing group) is claimed. The photosensitive material contains ≥ 1 of I. The material shows high sensitivity, and gives low fog images with good gradation and storage stability.

MSTR 1

G2---G1

G1 = 6

G2 = acylamino / SO2NH2 / CONH2 Patent location: claim 1

Note: additional ring formation also claimed

L5 ANSWER 100 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 127:42394 MARPAT

TITLE: Compound which changes the UV absorption with H+

concentration

INVENTOR(S): Jinbo, Yoshihiro; Nigorikawa, Kazunori; Waji, Naotaka

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE _____ JP 1995-252523 JP 09095669 Α 19970408 19950929 PRIORITY APPLN. INFO.: JP 1995-252523 19950929 GΙ

R9 R8 R10 _R5 R^{1} OR 0 R11 Ν R12 R6 R^2 ${\mathbb R}^4$ R3

AB The title compound, suited for use as a UV absorber and a recording material, is styryl quinoline derivs. represented by I [R0 = alkyl, aryl, and heterocyclic; R1-4 = H, halo, alkyl, aryl, cyano, etc.; R5-6 = H, and alkyl; and R7-12 = H, halo, aryl, cyano, etc.]. The increase in the H+ concentration of the solution transforms the quinoline form to the quinolinium form

in which the UV absorption spectra are dissimilar to quinoline from.

Ι

MSTR 1

= CONH2 (opt. substd.) / acylamino

Patent location: claim 2

Note: additional ring formation also claimed

ANSWER 101 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 127:5099 MARPAT

TITLE: Preparation of pyridazine derivatives for the treatment of endotoxin shock and kidney diseases

Ishida, Akihiko; Honma, Koichi; Tanifuji, Michihisa;

INVENTOR(S): Nishama, Nobusuke; Okumura, Fumikazu

PATENT ASSIGNEE(S): Tanabe Seiyaku Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 28 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 09071534	A	19970318	JP 1996-164798	19960625
PRIORITY APPLN. INFO.	:		JP 1995-159261	19950626
GI				

AB The title compds. I [R1 = alkyl, etc.; R2 = H, alkyl; X = C0, etc.; Alk = bond, alkylene; dotted line indicates optional double bond] are prepared When treated with the title compound II at 100 mg/kg orally, mice with endotoxin shock showed 90% survival.

MSTR 1

G3 = quinolinyl (opt. substd. by 1 or more G6)

G6 = loweralkylamino / CONH2

G11 = bond

Derivative: or pharmacologically acceptable salts

Patent location: claim 1

L5 ANSWER 102 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 127:5014 MARPAT

TITLE: Synthesis of substituted N-heteroaromatic compounds by

combinatorial chemistry

INVENTOR(S): Smith, Robert L.; Kumaravel, Gnanasambandam; Kuhla,

Donald E.

PATENT ASSIGNEE(S): Versicor, Inc., USA; Smith, Robert, L.; Kumaravel,

Gnanasambandam; Kuhla, Donald E.

SOURCE: PCT Int. Appl., 41 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE							APPLICATION NO. DATE										
 WO	9715	 557		 A	 1	 1997	0501		M.	 0 19	 96-U	 S171	 77	1996	 1025		
	W:	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CZ,	DE,	DK,
		EE,	ES,	FI,	GB,	GE,	HU,	IL,	IS,	JP,	ΚE,	KG,	KP,	KR,	KZ,	LC,	LK,
		LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,
		RU,	SD,	SE,	SG,	SI,	SK,	ΤJ,	TM,	TR,	TT,	UA,	UG,	US,	UZ,	VN	
	RW:	KΕ,	LS,	MW,	SD,	SZ,	UG,	ΑT,	BE,	CH,	DE,	DK,	ES,	FΙ,	FR,	GB,	GR,
		ΙE,	ΙT,	LU,	MC,	NL,	PT,	SE,	BF,	ΒJ,	CF,	CG,	CI				
US	5886	186		Α		1999	0323		U	S 19	95-5	4800	9	1995	1025		
AU	9675.	225		A		1997	0515		A	J 19	96-7	5225		1996	1025		
PRIORIT	Y APP	LN.	INFO	.:					U	S 19	95-5	4800	9	1995	1025		
									M	O 19	96-U	S171	77	1996	1025		

OTHER SOURCE(S): CASREACT 127:5014

GΙ

AB N-heteroarom. compds. I (W, X, Y, Z = bond, CR1; R1, R2 = H, halo, alkyl, alkenyl alkynyl, alkoxy, amino, acyl, CN, sulfhydryl, alkylthio, aryl, OH, carbamoyl, NO2, CF3, carbocycle), i.e. libraries of substituted N-heteroarom. compds., were prepared using polymer-supported reagents and featuring the reaction of O-linked heteroarom. N-oxides with nucleophiles to produce the substituted N-heteroarom. compds. Thus, II was prepd from 2-chloropyridine-N-oxide and N-(cyclohexen-1-yl)-morpholine using the acid chloride resin formed from the acyl chlorination of polyacrylic acid with SO2C12.

MSTR 2

G3----G2

G1 = 53 / 31

G3 = 92

G8 = heteroaryl G14 = 48

 $G19 = 115-90 \ 116-93$

Patent location: claim 12

Note: additional ring formation and substitution also

claimed

L5 ANSWER 103 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 126:199573 MARPAT

TITLE: Heterocyclylcarboxamide derivatives for use as

neurotransmitter agonists

Birch, Alan Martin; Heal, David John; Kerrigan, Frank; Martin, Keith Frank; Needham, Patricia Lesley; INVENTOR(S):

Sargent, Bruce Jeremy

PATENT ASSIGNEE(S): Knoll Aktiengesellschaft, Germany

PCT Int. Appl., 93 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	rent	NO.		KI	ND	DATE	TE APPLICATION NO.				DATE							
	9703 W:	071 AU,	BG,	A BR,	1 CA,	1997 CN,	0130 CZ,	GE,	WO HU,	0 19 IL,	96-E JP,	P289 KR,	0 LV,	1996 MX,	NO,	NΖ,		
														KZ,				
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	ĿΤ,	FR,	GB,	GR,	IE,	TT,	LU,	MC,	NL,	PT,	SE
CA	2223	4/2		A	1	199/	0130		C.	A 19	96-2	2234	72	1996	0/02			
AU	9665	172		A	_	1997	0210		А	U 19	96-6	5172		1996	0702			
		90																
EP	8391	.45		A	1	1998	0506		E.	P 19	96-9	2484	7	1996	0702			
EP		45																
	R:					DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	PT,	IE,	
		SI,	LV,	FI														
CN	1190	967		A		1998	0819		C1	N 19	96-1	9547	7	1996 1996 1996 1996	0702			
CN	1071	755		С		2001	0926											
BR	9609	506		A		1999	0601		B	R 19	96-9	506		1996	0702			
JP	1150	8599		Т		1999	0727		J:	P 19	96-5	0547	1	1996	0702			
HU	9901	485		A	2	2000	0728		H	U 19	99-1	485		1996	0702			
HU	9901	485 147		A	3	$\angle UUI$	U348											
RU	2169	147		С	2	2001	0620		R	U 19	98-1	0244	1	1996	0702			
IL	1225	40		A		2001	1031		I	L 19	96-1	2254	0	1996	0702			
AT	2535	73		Т		2003	1115		A'	Т 19	96-9	2484	7	1996	0702			
IN	1996	MA01	230	А		2005	0304		I	N 19	96-M	A123	0	1996 1996 1996	0711			
ZA	9605	921		А		1998	0112		\mathbf{z}	A 19	96-5	921		1996	0712			
TW	4540	06		В		2001	0911		T	w 19	96-8	5115	692	1996 1996	1219			
US	5935	973		А		1999	0810		IJ	S 19	98-9	8167	1	1998	0105			
														1998				
PRIORITY				.:					G:	В 19	95-1	4380		1995 1996	0713			

GI

$$\begin{array}{c|c} \text{C1} & \text{O} & \text{CH}_2\text{N} \\ \hline \\ \text{O} & \text{H}_2\text{N} \\ \end{array}$$

AB Title compds. I [A, B = CH2, O; R1 = optional substituent(s); R2-R4 = H, (un)substituted alkyl; U = (un)branched alkylene; Q = N-containing divalent group; T = heterocyclylcarbonyl attached to N in Q] were prepared for use in treating central nervous system disorders. Thus, the benzodioxane II was prepared from 5-chloro-2-hydroxybenzaldehyde, (R)-glycidyl tosylate, and 4-aminomethylpiperidine in 8 steps. II had a Ki for 5_HT1 α receptor binding of 41.5 nM and also bound to the α 2D, D2, and α 1 receptors.

MSTR 1

$$G7 = 393$$

$$G9 = 351$$

$$G23 = 130 / 136 / 198$$

G26 = alkyl < containing 1-5 C >

Derivative: and pharmaceutically acceptable salts

Patent location: claim 1

Note: substitution is restricted

L5 ANSWER 104 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 125:320547 MARPAT

TITLE: Synergistic fungicidal compositions made of quinoline

derivatives and cytochrome b/c inhibitors

INVENTOR(S): Koehle, Harald; Ammermann, Eberhard; Bayer, Herbert;

Wagner, Oliver; Roehl, Franz

PATENT ASSIGNEE(S): BASF A.-G., Germany SOURCE: PCT Int. Appl., 36 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	TENT NC		KI	ND	DATE APPLICATION NO.							Э.	. DATE				
WO	963201	5	. –– A	1	1996	1017		M	 J 19	 96-Е	P129	8	1996	0325			
	W: A	U, BG,	BR,	CA,	CN,	CZ,	HU,	JP,	KR,	MX,	NO,	NΖ,	PL,	SG,	SK,	TR,	
	Ü	A, US,	ΑM,	ΑZ,	BY,	KG,	KΖ,	MD,	RU,	ΤJ,	$_{ m IM}$						
	RW: A	T, BE,	CH,	DE,	DK,	ES,	FΙ,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE
CA	221551	4	A	1	1996	1017		C	A 19	96-2	2155	14	1996	0325			
AU	965148	6	A		1996	1030		A	J 19	96-5	1486		1996	0325			
EP	820232		A	1	1998	0128		E:	P 19	96-9	0813	1	1996	0325			
	R: A	T, BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	NL,	SE,	PT,	ΙE,	FΙ	
CN	118099	5	A		1998	0506		C1	N 19	96-1	9313	9	1996	0325			
HU	980163	0	A	2	1998	1130		H	J 19	98-1	630		1996	0325			
BR	960482	-											1996	0325			
JP	115034	35	T		1999	0326		J:	P 19	96-5	3067	2	1996	0325			
ZA	960270	9	A		1997	1006		$\mathbf{Z}_{\mathbf{z}}$	A 19	96-2	709		1996	0404			
PRIORITY	Y APPLN	. INFC	.:					D:	E 19	95-1	9513	404	1995	0408			
								M) 19	96-E	P129	8	1996	0325			

GΙ

The title fungicides comprise compds. that inhibit the respiration of cytochrome complex III and a quinoline derivative I (m = 1-6; R = H, cyano, nitro, hydroxy, mercapto, amino, carboxyl, aminocarbonyl, aminothiocarbonyl, sulfo, aminosulfonyl, halogen, alkyl, haydroxyalkyl, alkxoyalkyl, alkoxy, alkoxyalkoxy, alkylthio, alkylamino, dialkylamino, alkylsuphonyl, alkylsulfoxyl, alkylsulfonyloxy, alkylcarbonyl, alkylcarbonylamino, etc; R1 = H, cyano, nitro, hydroxy, mercapto, amino, carboxyl, aminocarbonyl, etc.).

MSTR 1

G1 = CONH2 / SO2NH2 / alkylamino <containing 1-6 C> (opt. substd. by 1 or more halo)

Patent location: claim 1

L5 ANSWER 105 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 125:114487 MARPAT

TITLE: CNS-Active pyridinylurea derivatives INVENTOR(S): Forbes, Ian Thomson; Jones, Graham Elgin

PATENT ASSIGNEE(S): Smithkline Beecham P.L.C., UK

SOURCE: PCT Int. Appl., 24 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO	. KIND	DATE	APPLICATION NO.	DATE
WO 961193	0 A1	19960425	WO 1995-EP3944	19951005
W: J:		r DK EG ED	GB, GR, IE, IT, LU	MC NI DT SE
EP 788499	A1		EP 1995-934135	
R: A			IT, LI, NL, SE	
JP 105085	84 T	19980825	JP 1995-512907	19951005
US 586658	6 A	19990202	US 1997-817580	19970417
PRIORITY APPLN	. INFO.:		GB 1994-20999	19941018
			WO 1995-EP3944	19951005

GΙ

$$Q1 = R6 \xrightarrow{R7} R7 R6$$

$$X \times R4 \qquad Q2 = R8 \times R4$$

$$Y \times R5 \qquad Y \times R5$$

$$W \times R4 \qquad X \times R4 \qquad X \times R4$$

$$Y \times R5 \qquad Y \times R5$$

The invention relates to heterocyclic compds. R1-G-N(R2)-CO-R3 [I; G = Ph AB ring, quinoline or isoquinoline nucleus, or a 5- or 6-membered aromatic heterocycle containing 1-3 heteroatoms (N, O, and/or S); R1 = H, alkyl, alkylthio, cyano, NO2, halo, CF3, amino, etc.; R2 = H, alkyl; R3 = group O1 or O2; X = Y = N, or one of X and Y = N and the other = C or CH; R4, R5 = alkyl, alkoxy, OH, halo, NO2, (un) substituted Ph, etc.; or R4R5 forms (un)substituted 5-membered carbo- or heterocyclic ring; R6, R7, R8 = H, alkyl]. Compds. I are 5-HT2C receptor antagonists, and some or all of them are also $5-\mathrm{HT2B}$ antagonists. They are useful in the treatment of a variety of CNS and GI disorders. For example, 5,6-dichloronicotinic acid underwent sulfurization in the 6-position by thiourea (87%) and S,O-dimethylation with MeI (50%) to give Me 3-chloro-2-(methylthio)pyridine-5-carboxylate. This was converted to the corresponding hydrazide (32%) and then the carbonyl azide (72%). latter was decomposed in refluxing PhMe, and the intermediate isocyanate treated with 3-aminopyridine, to give 85% title compound II. The three example compds. had pKi of 7.4-8.1 in a test for displacement of [3H]-mesulergine from rat or human 5-HT2C clones, expressed in 293 cells in vitro.

MSTR 1

Derivative: or salts Patent location: claim 1

Note: additional ring formation specified

L5 ANSWER 106 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 125:10629 MARPAT

TITLE: The alkoxylation of heterocyclic compounds in the

presence of fluorine

INVENTOR(S): Chambers, Richard Dickinson; Skinner, Christopher

John; Sandford, Graham

PATENT ASSIGNEE(S): Bnfl Fluorochemicals Ltd., UK

SOURCE: PCT Int. Appl., 16 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

W: JP, US

RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

ZA 9506176 A 19960308 ZA 1995-6176 19950725 PRIORITY APPLN. INFO.: GB 1994-14973 19940726

AB A method for introducing an alkoxy, acyloxy, alkenyloxy, aryloxy, etc., group into a nitrogen-containing heterocyclic aromatic compound is achieved in high

yield by reacting a compound containing the functionalizing group [e.g., an (un)substituted alc., acid, etc.] with the heterocyclic aromatic compound in the presence of fluorine. Thus, pyridine was reacted with EtOH in the presence of fluorine gas, producing 2-ethoxypyridine in 50% yield.

MSTR 2

G5-G1

G5 = 59

G8 = NH2

G9 = alkylcarbonylamino / CONH2 / 24

02S----G8

Patent location: disclosure

L5 ANSWER 107 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 124:316867 MARPAT

TITLE: Carbapenem derivatives containing a bicyclic

substituent

INVENTOR(S):
Arnould, Jean-Claude

PATENT ASSIGNEE(S): Zeneca Limited, UK; Zeneca-Pharma

SOURCE: Eur. Pat. Appl., 27 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 695753	A1	19960207	EP 1995-305428	19950803
R: AT, BE,	CH, DE	, DK, ES, FR,	GB, GR, IE, IT, LI	, LU, MC, NL, PT, SE
US 5607928	А	19970304	US 1995-508698	19950728
CA 2155493	A1	19960206	CA 1995-2155493	19950804
CA 2155493	С	20070501		
JP 08059664	A	19960305	JP 1995-201126	19950807
JP 4031538	В2	20080109		
PRIORITY APPLN. INFO	.:		EP 1994-401814	19940805
GI				

$$R^{1}$$
 R^{2}
 $CH_{2}XR$
 Me
 $CH_{2}O$
 $CO_{2}H$ I
 $CO_{2}Na$
 $CO_{2}Na$

AB Bactericidal (no data) carbapenems I [R = aryl, heteroaryl; R1 = CH2OH, CHMeOH, CHMeF; R2 = H, C1-4 alkyl; X = O, S] and pharmaceutically acceptable salts or in vivo hydrolyzable esters thereof, were prepared Thus, (3S, 4R, 1'R, 1''R) - 1 - (allyloxycarbonyltriphenylphosphoranylidenemethyl) -3 - (1 - hydroxyethyl) -4 - [1 - (hydroxymethylcarbonyl)ethyl]azetidin -2 - one was treated with 5 - hydroxy -1 - tetralone, followed by ester hydrolysis to give the carbapenem II.

MSTR 1A

= CONH2 / SO2NH2 / 141

G6

G12-SO₂-G13

G12 = NH

Derivative: and pharmaceutically acceptable salts

Derivative: or protected derivatives

Patent location: claim 1

Note: substitution is restricted Note: also incorporates claim 16

L5 ANSWER 108 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 124:146140 MARPAT

TITLE: Preparation of N-(3- and 5-

isoxazolyl)biphenylsulfonamides as endothelin receptor

li1gands

INVENTOR(S): Chan, Ming F.; Raju, Bore G.; Castillo, Rosario S.;

Kois, Adam; Wu, Chengde; Balaji, Vitukudi

PATENT ASSIGNEE(S): ImmunoPharmaceutics, Inc., USA

SOURCE: U.S., 17 pp. Cont.-in-part of U.S. Ser. No. 100, 565,

abandoned.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 10

PATENT INFORMATION:

PATENT NO. KI		APPLICATION NO.	DATE
US 5464853 A	19951107	US 1993-142159	19931021
US 5591761 A	19970107	US 1994-222287	19940405
CA 2161346 A	1 19941208	CA 1994-2161346	19940520
CA 2161346 C			
WO 9427979 A	1 19941208	WO 1994-US5755	19940520
W: AT, AU, BB,	BG, BR, BY, C	CA, CH, CN, CZ, DE, DK,	ES, FI, GB, GE,
HU, JP, KG,	KP, KR, KZ, L	K, LU, LV, MD, MG, MN,	MW, NL, NO, NZ,
PL, PT, RO,	RU, SD, SE, S	SI, SK, TJ, TT, UA, US	
RW: AT, BE, CH,	DE, DK, ES, F	R, GB, GR, IE, IT, LU,	MC, NL, PT, SE,
BF, BJ, CF,	CG, CI, CM, G	GA, GN, ML, MR, NE, SN,	TD, TG
AU 9469646 A	19941220	AU 1994-69646	19940520
AU 691813 B	2 19980528		
GB 2285625 A	19950719	GB 1995-3693	19940520
GB 2285625 B	19971210		
EP 699191 A	1 19960306	EP 1994-918081	19940520
EP 699191 B	1 19981216		
		R, GR, IE, IT, LI, LU,	
US 5571821 A	19961105	US 1994-247072 JP 1995-500856	19940520
JP 08510744 T	19961112	JP 1995-500856	19940520
		EP 1998-109339	
R: AT, BE, CH,	DE, DK, ES, F	R, GR, IT, LI, LU, NL,	SE, MC, PT, IE
AT 174592 T	19990115	AT 1994-918081	19940520
ES 2127397 T	3 19990416	ES 1994-918081	19940520
RU 2151144 C	1 20000620	RU 1995-121744	19940520
		EP 2000-119107	
EP 1069114 A	3 20010131		
R: AT, BE, CH,	DE, DK, ES, F	R, GR, IT, LI, LU, NL,	SE, MC, PT, IE

US 5594021	А	19970114	US	1995-477223	19950606
US 5962490	A	19991005	US	1996-721183	19960927
US 6030991	A	20000229	US	1996-730633	19961206
AU 9860585	A	19980604	AU	1998-60585	19980331
AU 724575	В2	20000928			
US 6331637	В1	20011218	US	1999-274280	19990322
AU 9935803	А	19990916	AU	1999-35803	19990622
AU 726595	В2	20001116			
US 2001003695	8 A1	20011101	US	2000-749716	20001227
US 6541498	В2	20030401			
PRIORITY APPLN. IN	FO.:		US	1993-65202	19930520
			US	1993-100125	19930730
			US	1993-100565	19930730
			US	1987-100865	19870925
			US	1990-416199	19900515
			US	1993-142159	19931021
			US	1993-142552	19931021
			US	1993-142631	19931021
			US	1994-222287	19940405
				1994-918081	19940520
			EP	1998-109339	19940520
				1994-247072	19940520
				1994-US5755	19940520
			US	1995-416199	19950404
			US	1995-417075	19950404
			US	1995-477223	19950606
			AU	1996-55367	19960404
			WO	1996-US4759	19960404
				1996-721183	19960927
				1996-730633	19961206
			US	1999-439802	19991112
AD DOCCOMIID1 LT.	D1 /	A 1 1 1 1	7	DO - 111	/ \ l + - + +

AB R2SO2NHR1 [I; R1 = (un)substituted aryl; R2 = alkenyl, (un)substituted phenyl(alkyl), (un)substituted PhCH:CH, etc.] were prepared Thus, 5-amino-3,4-dimethylisoxazole was amidated by 4-PhC6H4SO2Cl to give N-(3,4-dimethyl-5-isoxazolyl)-4-biphenylsulfonamide;. I had IC50 of <100 μ M against ligand binding at endothelin ETA and ETB receptors in vitro.

MSTR 3

$$G3 = 55$$

G4 = NHOH / CONH2 (opt. substd.)
Patent location: disclosure

Note: substitution is restricted

Note: additional ring formation allowed

L5 ANSWER 109 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 123:49819 MARPAT

TITLE: Marine antifouling coating.

INVENTOR(S): Anthoni, Uffe; Christophersen, Carsten; Nielsen, Per

Halfdan; Kjaer, Eva Bie; Musaeus, Gruska Folkmann;

Schultz, Ann Christina

PATENT ASSIGNEE(S): J.C. Hempel's Skibsfarve-Fabrik A/S, Den.

SOURCE: PCT Int. Appl., 53 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

P.F	ATENT	NO.		KI	KIND DATE					APPLICATION NO.					DATE			
WC	9511	592		A.	1	1995	0504		Mo	O 19	94-D	K405		1994	1028			
	W:	ΑM,	ΑT,	ΑU,	BB,	BG,	BR,	BY,	CA,	CN,	CZ,	CZ,	DE,	DK,	EE,	ES,	FΙ,	
		FΙ,	GE,	HU,	JP,	KG,	KP,	KR,	KΖ,	LK,	LR,	LT,	LV,	MD,	MG,	MN,	NO,	
		NΖ,	PL,	RO,	RU,	SI,	SK,	SK,	ТJ,	TT,	UA,	US,	UΖ,	VN				
	RW:	ΚE,	MW,	SD,	SZ,	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	IT,	LU,	
		MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,	MR,	NE,	SN,	
		TD,	ΤG															
JA	J 9480	576		Α		1995	0522		A	J 19	94-8	0576		1994	1028			
EI	7255	63		A.	1	1996	0814		E	P 19	94 - 9	3151	9	1994	1028			
	R:	BE,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	ΙΤ,	NL,	PΤ						
PRIORIT	Y APP	LN.	INFO	. :					DI	K 19	93-1	226		1993	1029			
									M	O 19	94-D	K405		1994	1028			

$$R^4$$
 R^3 R^2 R^3 R^4 R^3 R^2 R^3 R^4 R^3 R^4 R^3 R^4 R^3 R^2 R^4 R^3 R^4 R^3 R^4 R^3 R^4 R^3 R^4 R^3 R^4 R^4 R^3 R^4 R^4

AB The title coating comprises a quinoline compound I [R1,R2,R4,R5,R6,R7 = H,OH,(un)substituted alkyl, etc.;R3 = R1,(un)substituted 1-azabicyclo[2,2,2]octylalkyl] or an N-oxide or a salt thereof. I exhibited activity against Enteromorpha, Amphora, Nitocra, and Balanus.

MSTR 1

GΙ

G1 = CONH2 / alkylaminosulfonyl <containing 1-12 C>

(opt. substd.)

G6 = alkylamino <containing 1-12 C> (opt. substd.)

G9 = N

Derivative: or salts Patent location: claim 1

L5 ANSWER 110 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 122:303102 MARPAT

TITLE: Photothermographic materials. INVENTOR(S): Kirk, Mark P.; Mott, Andrew W.

PATENT ASSIGNEE(S): Minnesota Mining and Manufacturing Co., USA

SOURCE: Eur. Pat. Appl., 14 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 631176	A1	19941228	EP 1994-304069	19940607
EP 631176	B1	20001213		
R: BE, DE,	FR, GB	, IT, NL		
US 5460938	A	19951024	US 1994-247651	19940523
CA 2124755	A1	19941209	CA 1994-2124755	19940531
JP 07002781	A	19950106	JP 1994-125023	19940607
JP 2801856	В2	19980921		
US 5594143	A	19970114	US 1995-464162	19950605
PRIORITY APPLN. INFO	.:		GB 1993-11790	19930608
			US 1994-247651	19940523

GI

AB A compound having a nucleus of the formula I are suitable for use as image stabilizers and anti-fog agents in photothermog. materials and exhibit acceptably low sensitization of human skin.

MSTR 1

G1 = CONH2 (opt. substd.) / 22 / SO2NH2

HN----C(O)-R

Patent location: claim 2

L5 ANSWER 111 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 121:311780 MARPAT

TITLE: Silver halide color photographic light-sensitive

material.

INVENTOR(S): Ueda, Fumitaka; Nishigaki, Junji PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 76 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATI	ENT NO.		KIND	DATE	API	PLICATION N	O. DATE
EP (600518		A2	19940608	EP	1993-11955	6 19931203
EP (600518		A3	19950329			
EP (600518		B1	19980325			
	R: BE	, DE,	FR, GH	B, NL			
JP (06175289	9	A	19940624	JP	1992-34999	8 19921203
US !	5449594		A	19950912	US	1993-15974	8 19931201
PRIORITY	APPLN.	INFO.	. :		JP	1992-34999	8 19921203
GI							

A Ag halide color photog. light-sensitive material includes a support AΒ having provided thereon at least 1 blue-sensitive Ag halide emulsion layer, at least 1 green-sensitive Ag halide emulsion layer, at least 1 red-sensitive halide emulsion layer, and at least 1 hydrophilic colloid layer. The hydrophilic colloid layer contains a compound represented by I, a Ag halide emulsion layer having an interlayer effect on the red-sensitive layer is also provided, and the layer with the interlayer effect contains a Aq halide emulsion spectrally sensitized with a sensitizing dye II or III. : In I R represents a hydrogen atom, alkyl. alkenyl, aryl, a heterocyclic ring, ureido, sulfonamide, sulfamoyl, sulfonyl, sulfinyl, alkylthio, arylthio, oxycarbonyl, acyl, carbamoyl, cyano, alkoxy, aryloxy, amino, or amide; Q represents -O- or -NR2- wherein R2 represents a hydrogen atom, alkyl, aryl, or a heterocyclic group; R3, R4, and R5 each represent a hydrogen atom, alkyl, or aryl, and R4 and R5 being able to be bonded to each other to form a 6 membered ring; R6 represents a hydrogen atom, alkyl, aryl, or amino; L1, L2, and L3 each represent methine, and k is an integer of 0 or 1. In II R11, R12, R13, and R14 may be the same or different and each represent a hydrogen atom, a halogen atom, alkyl, aryl, alkoxy, aryloxy, aryloxycarbonyl, alkoxycarbonyl, amino, acyl, cyano, carbamoyl, sulfamoyl, carboxyl, or an acyloxy group, R11 and R12 or R13 and R14 not representing a hydrogen atom simultaneously; R15 and R16 may be the same or different and each represent an alkyl group; R17 represents an alkyl having not less than three carbon atoms, aryl, or aralkyl group; X represents a counter anion, and m is an integer of 0 or 1, and m = 0 when intramol. salt is to be formed. In III R21, R22, R23, R24, R25, R26, R27, R28, R29, and R30 each have the same meaning as that of R11, R31 and R32 each have the same meaning as that of R15; Y represents a sulfur atom, a selenium atom, or an

oxygen atom; X has the same meaning as that of X1; and n has the same meaning as that of m. The material provides good coloration and has high speed and high graininess.

MSTR 3

G1 = CONH2 (opt. substd.) / SO2NH2 (opt. substd.) / acylamino

Patent location: claim 1

L5 ANSWER 112 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 121:289519 MARPAT

TITLE: Silver halide photographic material

INVENTOR(S): Kato, Takashi; Ikeda, Tadashi
PATENT ASSIGNEE(S): Fuji Photo Film Co Ltd, Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 24 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 06102614	A	19940415	JP 1992-254565	19920924
US 5462851	Α	19951031	US 1993-121740	19930916
PRIORITY APPLN. INFO.	:		JP 1992-254565	19920924
GI				

$$V^{4}$$
 V^{5}
 V^{1}
 V^{1}
 V^{2}
 V^{1}
 V^{2}
 V^{1}
 V^{2}
 V^{3}
 V^{2}
 V^{1}
 V^{2}
 V^{3}
 V^{2}
 V^{1}
 V^{2}
 V^{3}
 V^{3}
 V^{2}
 V^{3}
 V^{3

Ι

$$V^{9}$$
 V^{8} V^{7} V^{7} V^{7} V^{7} V^{10} V^{10}

AB The title photog. material contains ≥ 1 compound selected from I and II [Z1,2 = 5- or 6-membered N-containing heterocyclic ring; R1-5 = alkyl; R3,6 = alkyl, aryl, heterocyclyl; V1-12 = H, substituent; L1-10 = methine; M1,2 = counter ion; m1,2 \geq 0; n1,2 = 0, 1]. This material shows sharp absorption in the IR region.

MSTR 1

G13 = 55 / acylamino

G15-G14

G14 = NH2 / morpholino

G15 = C(0) / S02

Patent location: claim 1

L5 ANSWER 113 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 121:205225 MARPAT

TITLE: Quinoline-derivative leukotriene antagonists

INVENTOR(S):
Daines, Robert A.; Pendrak, Israil

PATENT ASSIGNEE(S): SmithKline Beecham Corp., USA

SOURCE: PCT Int. Appl., 17 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

W: JP, US

RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE PRIORITY APPLN. INFO.:

US 1992-996220 19921223
GI

AB The title compds. [I; A = CH2, CH0H, CO, (un)substituted NH, O, etc.; R = (un)substituted C1-20 aliphatic; R1 = 5-tetrazolyl, CO2H, (un)substituted aminoalkyl, etc.; R2 = H, halogen, CF3, CN, lower alkyl, lower alkyloxy, etc.; R3 = H, halogen, lower alkyl, C1-6 acyl; Z = (un)substituted NH, S(0)q, CO; q = 0-2], useful as leukotriene antagonists (no data), especially

LTB4 (no data), are prepared and I-containing formulation presented. Thus, $7-[1-\text{thia}-2-[2-(E-2-\text{carboxyethenyl})-3-[8-(4-\text{methoxyphenyl})\,\text{octyloxy}]-6-$ pyridyl]ethyl]quinoline Li salt was prepared from 7-hydroxyquinoline in 5 steps.

MSTR 1

for

G18 = quinolinyl (opt. substd. by (1-2) G19)

G19 = 66 / CONH2 (opt. substd.)

G21—G22

G21 = NH

G22 = cycloalkyl <containing 4-10 C>

Derivative: or pharmaceutically acceptable salts or N-oxides

Patent location: claim 1

L5 ANSWER 114 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 121:205125 MARPAT

TITLE: Preparation of [[(carboxyheterocyclyl)carbamoyl]pyrrol

idinylthio]carbapenems as antibiotics

INVENTOR(S): Jung, Frederic Henri; Arnould, Jean Claude

PATENT ASSIGNEE(S): Zeneca Ltd., UK; Zeneca Pharma S.A.

SOURCE: Eur. Pat. Appl., 27 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 581500	A1	19940202	EP 1993-305607	19930716
EP 581500	B1	19980909		
R: AT, B	E, CH, DE	, DK, ES, FR,	GB, GR, IE, IT, LI	, LU, MC, NL, PT, SE
CA 2099818	A1	19940122	CA 1993-2099818	19930705
AT 170859	T	19980915	AT 1993-305607	19930716
ES 2121585	Т3	19981201	ES 1993-305607	19930716
JP 06179674	A	19940628	JP 1993-177903	19930719
US 5441949	A	19950815	US 1994-307048	19940916
PRIORITY APPLN. IN	FO.:		EP 1992-402105	19920721
			US 1993-86836	19930707

GΙ

$$\begin{array}{c|c} & H & S & CONR^3ZCO_2H \\ \hline & N & NR^2 & \end{array}$$

AB Title compds. [I; R1 = MeCH(OH), MeCHF, CH2OH; R2,R3 = H, alkyl; Z = (iso)quinolinediyl, quinazolinediyl, quinoxalinediyl, etc.] were prepared Thus, disodium (1R,5S,6S,8R,2'S,4'S)-2-[2-(8-carboxyquinol-6-ylcarbamoyl)pyrrolidin-4-ylthio]-6-(1-hydroxyethyl)-1-methylcarbapenem-3-carboxylate, prepared in 5 steps from 6-amino-8-carboxyquinoline (preparation given), had MIC of 0.13 and 0.03 μ g/mL against Staphylococcus aureus Oxford and Escherichia coli DCO, resp.

Ι

MSTR 1

$$G1$$
 H
 $G2$
 $C(0) \cdot G3 - G5$
 CO_2H

G3 = NH

G5 = quinolinyl (substd. by (1-4) G10)

G10 = CONH2

Derivative: or pharmaceutically acceptable salts or in-vivo

hydrolysable esters; or protected derivatives

Patent location: claim 1

Note: also incorporates claim 8

L5 ANSWER 115 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 121:167055 MARPAT

TITLE: Photothermographic imaging materials and antifoggants

therefor.

INVENTOR(S): Oliff, David B.; Kirk, Mark P.

PATENT ASSIGNEE(S): Minnesota Mining and Manufacturing Co., USA

SOURCE: Eur. Pat. Appl., 21 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 600587	A1	19940608	EP 1993-307740	19930929
EP 600587	B1	19960214		
R: DE, FR,	GB, IT			
US 5939248	A	19990817	US 1993-126331	19930924
JP 06202268	A	19940722	JP 1993-252998	19931008
PRIORITY APPLN. INFO.	.:		GB 1992-21383	19921012
GI				

AB A photothermog. material having a photosensitive medium comprising: photosensitive Ag halide, a reducible Ag source, a reducing agent for Ag ion, a hydrobromic acid salt of a N-containing heterocyclic ring or fused ring nucleus associated with a pair of bromine atoms characterized in that the photosensitive medium addnl. comprises as antifoggant, substantially in the absence of an antifoggant amount of Hg and other heavy metal salts, a tribromomethyl ketone compound of a general formula I (R = alkyl, aryl, a carbocyclic ring or fused ring nucleus, heterocyclic ring or fused ring

nucleus).

MSTR 2

G1 H—Br Br—Br

G1 = 24

G5 G5 G5 G5 G5 24 G5

G5 = acylamino / SO2NH2 / CONH2 Patent location: claim 7

Note: substitution is restricted

L5 ANSWER 116 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 121:108803 MARPAT

TITLE: Preparation of tetrazoles as intermediates for

photographic couplers

INVENTOR(S): Ookawa, Atsuhiro

PATENT ASSIGNEE(S): Fuji Photo Film Co Ltd, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 27 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	AP:	PLICATION NO.	DATE
JP 05331145	A	19931214	JP	1992-132707	19920525
JP 2881356	В2	19990412			
US 5362877	A	19941108	US	1993-64990	19930524
PRIORITY APPLN. INFO.	:		JP	1992-132707	19920525
OTHER SOURCE(S):	CA	SREACT 121:1088	303		
GI					

R10CONR²CN
$$R^3$$
 R^3 R^3

The title compds. I [R1 = alkyl, aryl, heterocyclic ring; R2 = alkyl, AB aryl; R3, R4 = H, alkyl, etc.; X2 = non-metallic atoms for forming 5- or 6-membered N-containing heterocyclic ring] were prepared by condensation of the appropriate amines with aldehydes (or ketones) and mercaptoheterocycles in the presence of a Lewis acid and/or a metal salt. Reaction of amine II (T = H) with paraformaldehyde and mercaptotetrazole III in the presence of BF3.OEt2 gave, after workup, 61% II (T = Q1), vs. 0% yield in the absence of Lewis acid or of metal salt.

MSTR 3

HS-G1

= 113 G1

= CONH2 (opt. substd.) / SO2NH2 (opt. substd.) / acylamino

Patent location: claim 1

ANSWER 117 OF 131 MARPAT COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 121:9027 MARPAT

TITLE: Preparation of (pyridiniomethyl)cephemcarboxylates and analogs as antibacterial agents

INVENTOR(S): Takamura, Norio; Saito, Kunio; Matsushita, Tadahiro;

Yamaguchi, Totaro

PATENT ASSIGNEE(S): Tanabe Seiyaku Co, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 47 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 05202062	A	19930810	JP 1992-53045	19920127
PRIORITY APPLN. INFO.	:		JP 1992-53045	19920127
GI				

AB The title compds. I [R1 = (protected) amino; R2 = (protected) OH, alkoxy; R3 = (protected) carboxyl; R4 = H, alkyl, CH2R41, etc.; R41 = nucleophilic moiety; R5 = (protected) carboxyl, CO2-; R6 = H, alkyl; the dotted line represents either a double bond or a single bond] were prepared Reaction of 7β -[(Z)-2-(2-aminothiazol-4-yl)-2-[(8-hydroxy-2-oxo-1H-quinoline-5-yl) (carboxyl)methyloxyimino]acetamido]cephalosporanic acid di-Na salt with pyridine in the presence of NaI gave cephem (Z)-II isolated as α and β isomers. The title compds. in vitro exhibited MIC values of 0.1-0.78 μ g/mL (against Staphylococcus aureus 209P JC-1) and MIC values

ΙI

Ι

of 0.78-1.56 $\mu g/mL$ against Pseudomonas aeruginosa Number 12.

MSTR 1

G8 = 71

G9 = 98

G16 = CONH2 / NHCHO

Derivative: or pharmacologically acceptable salts

Patent location: claim 1

L5 ANSWER 118 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 121:4152 MARPAT

TITLE: Metal complexes of hydroxyaryl-containing amino

carboxylic acid chelating agents

INVENTOR(S): Smith, Suzanne Virginia; Lambrecht, Richard Merle;

Schmidt, Peter Frederick; Lee, Fook Thean

PATENT ASSIGNEE(S): Australian Nuclear Science and Technology

Organisation, Australia SOURCE: Eur. Pat. Appl., 24 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PAT	TENT NO.		KIND	DATE		APPLICATION NO. DATE	
			A2 A3	19940406 19940824		EP 1993-305992 19930729	
	590766		В1	20000202			
	R: AT,					, GB, GR, IE, IT, LI, LU, NL, PT, SE	
	955066					EP 1999-112338 19930729	
EP	955066		А3	20020828			
	R: AT,	BE,	CH, DE	, DK, ES,	FR,	, GB, GR, IT, LI, LU, NL, SE, PT, IE	
AT	189396					AT 1993-305992 19930729	
PT	590766		T	20000731		PT 1993-305992 19930729	
ES	2146217		Т3	20000801		ES 1993-305992 19930729	
AU	9344374		А	19940203		AU 1993-44374 19930730	
AU	671465		В2	19960829			
JP	07285888	}	А	19951031		JP 1993-208689 19930731	
GR	3033352		Т3	20000929		GR 2000-401036 20000502	
PRIORITY	APPLN.	INFO	. :			AU 1992-3883 19920731	
						EP 1993-305992 19930729	

AB Complexes of a radioactive metal (especially 99mTc, 188Re, 186Re) with EDTA analogs XNHC(O)CH2N(CH2CO2H)[(CH2)kN(CH2CO2H)].scriptl.CH2C(O)NHY [I; X, Y = aryl or heteroaryl, especially (substituted) Ph, naphthyl, pyridyl, quinolinyl; k = 2-5; .scriptl. = 1-5] are prepared for use as imaging agents, e.g. to assess hepatobiliary function, or in radiolabeling of monoclonal antibodies, proteins, peptides, oligonucleotides, etc. for in vivo imaging or therapy. Thus, 2-amino-4-nitrophenol reacted with EDTA anhydride to produce I (X = Y = 2-hydroxy-5-nitrophenyl; k = 2; .scriptl. = 1) (II). The 99mTc complex of II, injected into rats, localized predominantly in the kidneys and somewhat less in the liver.

MSTR 1

G3 = quinolinyl (opt. substd. by 1 or more G7)

G7 = CONH2

Derivative: or pharmaceutically acceptable salts or complexes

with G10

Patent location: claim 1

L5 ANSWER 119 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 120:334755 MARPAT

TITLE: Color developer composition and photographic

processing using same

INVENTOR(S): Fujimoto, Hiroshi; Morimoto, Kyoshi; Furusawa,

Genichi; Myashita, Yosuke; Taniguchi, Masato

PATENT ASSIGNEE(S): Fuji Photo Film Co Ltd, Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 15 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 05188551 A 19930730 JP 1992-170973 19920629
PRIORITY APPLN. INFO.: JP 1991-197297 19910712
GI

AB The title color developer composition contains as additive ≥ 1 I [R1-4 = H, alkyl, aryl, aralkyl, halo, OH, NH2, alkoxy, COOH, SO3H, PO(OH)2, NO2, CN, heterocyclyl, carbamoyl, sulfamoyl, acyl, acylamino, alkylsulfonyl amino, arylsulfonyl amino, alkoxycarbonyl, ureido; R1,R2 may join to form a ring; m,n = 0-3]. Precipitation of the components of the above composition does not

occur during processing, the volume of the processing waste solution is reduced, and the developer solution is stable.

MSTR 1

$$G1 = 9-4 \ 10-2$$

G2 = SO2NH2 (opt. substd.)
G6 = CONH2 (opt. substd.) / acylamino

Patent location: claim 1

L5 ANSWER 120 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 120:191707 MARPAT

TITLE: 2-Substituted saccharin derivative proteolytic enzyme

inhibitors

INVENTOR(S): Hlasta, Dennis John; Desai, Ranjit Chimanlal;

Subramanyam, Chakrapani; Lodge, Eric Piatt; Dunlap, Richard Paul; Boaz, Neil Warren; Mura, Albert Joseph;

Latimer, Lee Hamilton

PATENT ASSIGNEE(S): Sterling Winthrop Inc., USA

SOURCE: Eur. Pat. Appl., 77 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 7

PATENT INFORMATION:

PA	TENT NO.	ř	IND	DATE	API	PLICATION NO).	DATE			
						1992-203469 GR, IE, IT,			NL,	PT,	SE
						1991-793033				ŕ	
						1992-25340					
AU	654581			19941110							
CA	2079822			19930516	CA	1992-207982	22	19921005			
				19930518	NO	1992-4401		19921113			
	303119			19980602							
HU	66873		A2	19950130	HU	1992-3566		19921113			
	103748					1992-103748		19921113			
	2101281			19980110	RU	1992-4381		19921113			
	05194444					1992-305295					
	5371074					1993-67637					
US	5650422		A	19970722	US	1994-270964	1	19940705			
US	5596012		A	19970121	US	1995-449152	2	19950524			
			A	19990223		1997-803297					
PRIORIT	Y APPLN. I	INFO.:				1991-793033		19911115			
						1989-347125		19890504			
						1989-347126		19890504			
						1990-514920		19900426			
						1993-67637					
					US	1994-270964	1	19940705			

GΙ

$$R^{4}$$

N (CH=CH) mC (R²) HL_nR¹

S
O
O

AB The title compds. I [L = 0, S, SO, SO2; R1 = (un)substituted Ph, (un) substituted heterocyclyl, etc.; R2 = H, lower alkoxycarbonyl, Ph, PhS; R3 = H, halogen, (un) substituted alkyl, Ph, lower alkoxy, lower alkoxycarbonyl, CN, etc.; R4 = H or 1-3 substituents selected from halogen, CN, NO2, NH2, etc.; m, n = 0, 1; when m = 0 then R1 can only be heterocyclyl and CHR2 can only be bonded to a ring N of R1; when m = 0, n = 1 and L is O, S, or SO, then R2-R4 = H; when m = 0, n = 1, L is S, R2, R4 = H and R3 = halogen; when m = 0, n = 1, and L is SO or SO2 then R2 is lower alkoxycarbonyl and R3 = R4 = H while $R1 \neq substituted Ph],$ useful for the treatment of degenerative diseases (no data), are prepared Thus, 2-hydroxymethyl-4-chlorosaccharin was reacted with thionyl chloride, producing 2-chloromethyl-4-chlorosaccharin (II). II demonstrated inhibition constant for human leukocyte elastase (rate of reactivation of enzyme to rate of inactivation of enzyme) of 0.5 nM and 26 nM for α -chymotrypsin.

MSTR 1A

G2 = bond G3 = 163

G12 G12 G12 G12 G12 163 N G12

G12 = alkylamino <containing 1-10 C> / CONH2 /

dialkylaminosulfonyl <each alkyl containing 1-10 C>

Patent location: claim 1

Note: substitution is restricted

L5 ANSWER 121 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 120:120563 MARPAT

TITLE: Method for processing silver halide color photographic

material

INVENTOR(S): Fujimoto, Hiroshi; Morimoto, Kyoshi; Furusawa,

Genichi; Myashita, Yosuke

PATENT ASSIGNEE(S): Fuji Photo Film Co Ltd, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 31 pp.

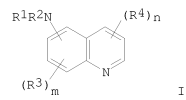
CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 05027394	A	19930205	JP 1991-202258	19910718
PRIORITY APPLN. INFO.	:		JP 1991-202258	19910718
GI				



AB In the title method involving the color development, desilvering, and washing or stabilization of a silver halide photog. material, the color developing solution contains one or more compds. represented by I. For I, R1-R4=H, alkyl, aryl, hydroxy, etc., R1 and R2 may together from a ring; m, n=0 to 3. The amount of replenishing solution for washing or stabilizing the photog. material is 3 to 50 times that of the amount of liquid brought from the preceding bath. The title method is economical and gives stable images.

MSTR 1

G3 = CONH2 (opt. substd.) / acylamino

G4 = SO2NH2 (opt. substd.)

Patent location: claim 1

Note: substitution is restricted

Note: additional ring formation possible

L5 ANSWER 122 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 120:120562 MARPAT

TITLE: Method for processing silver halide color photographic

material

INVENTOR(S): Furusawa, Genichi; Myashita, Yosuke; Fujimoto,

Hiroshi; Morimoto, Kyoshi

PATENT ASSIGNEE(S): Fuji Photo Film Co Ltd, Japan SOURCE: Jpn. Kokai Tokkyo Koho, 36 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 05027395	A	19930205	JP 1991-203633	19910719
PRIORITY APPLN. INFO.	:		JP 1991-203633	19910719
GI				

$$\mathbb{R}^{1}\mathbb{R}^{2}\mathbb{N}$$
 $\mathbb{R}^{4}\mathbb{N}$ $\mathbb{R}^{4}\mathbb{N}$ $\mathbb{R}^{3}\mathbb{N}$ $\mathbb{R}^{3}\mathbb{N}$ $\mathbb{R}^{3}\mathbb{N}$

AB The title method involves the treatment of the title material with a color developing solution containing a hydroxylamine derivative and a quinoline derivative

represented by I. For I, R1-R4 = H, alkyl, aryl, etc.; or R1 and R2 may together form a ring; m, n = 0 to 3. The title method is economical.

MSTR 1

G3 = CONH2 (opt. substd.) / acylamino

G4 = SO2NH2 (opt. substd.)

Patent location: claim 1

Note: substitution is restricted

Note: additional ring formation possible

L5 ANSWER 123 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 119:213908 MARPAT

TITLE: Silver halide photographic material

INVENTOR(S): Fukuwa, Junichi; Kobayashi, Akira; Goto, Kenji

PATENT ASSIGNEE(S): Konica Co., Japan

SOURCE: Can. Pat. Appl., 71 pp.

CODEN: CPXXEB

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CA 2065106	A1	19921005	CA 1992-2065106	19920403
JP 05197057	A	19930806	JP 1992-110787	19920403
PRIORITY APPLN. INFO.	. :		JP 1991-99626	19910404
GI				

AΒ A Ag halide photog. material for high-contrast dot image formation is disclosed. The material comprises a support and provided thereon a Ag halide emulsion layer and layers adjacent to the emulsion layer. The emulsion is subjected to desalinization comprising using denatured gelatin in the process of preparation thereof. At least one of the layers contains a hydrazine derivative and a compound selected from the group consisting of those represented by formulas A(CH2) nSC(:N+HR1) NHR1 X-(A=OH, SO3-, or N(R2)2;R1 = H, (substituted) alkyl having 1-5 C atoms, or (substituted) Ph; R2 =(substituted) alkyl having 1-5 C atoms; X-= an anion), (R3)2N(CH2)nSC(S)N(R4)2 (R3 = H, (substituted) alkyl having 1-5 C atoms, or (substituted) aryl; R4 = (substituted) alkyl having 1-5 C atoms or (substituted) Ph; n = an integer of 2-5), or I (Q = a group of atoms necessary to form a 5- or 6-membered heterocyclic ring which may be condensed with a benzene or heterocyclic ring; M = H, an alkali metal atom, an ammonium group, or an amine residue).

MSTR 3B

G1---G2

$$G1 = 232$$

$$G7 = 35 / CONH2 / 37$$

G14 = morpholino G15 = Ph INVENTOR(S):

Patent location: claim 1

L5 ANSWER 124 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 119:139102 MARPAT

TITLE: Antiproliferative derivatives of 4H-naphthol[1,2-

b]pyran and process for their preparation Dell, Colin Peter; Smith, Colin William

PATENT ASSIGNEE(S): Lilly Industries Ltd., UK SOURCE: Eur. Pat. Appl., 15 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE		PLICATION NO.	DATE	
	A1 B1			1992-309169	19921008	
			FR. GB.	GR, IE, IT, L	T. I.U. NI.	PT. SE
				1992-2079428		•
	A			1992-26216		
	B2	19950330		1331 10110		
	B6			1992-3035	19921005	
	А			1992-103356		
	C1			1992-5052861		
	A			1992-7717		
	В1	19991101	KR	1992-18309	19921007	
	A			1992-3910		
NO 301587	В1	19971117				
HU 62281	A2	19930428	HU	1992-3183	19921008	
HU 218916	A2 B	20001228				
CN 1073437	A	19930623	CN	1992-111625	19921008	
CN 1034938	С	19970521				
JP 0519447	7 A	19930803	JP	1992-269002	19921008	
AT 167859	7 A T	19980715		1992-309169		
ES 2117035	Т3		ES	1992-309169	19921008	
PRIORITY APPLN.	INFO.:		GB	1991-21358	19911009	
			GB	1992-13058	19920619	
GI						

AB The title compds. I [R1 = halogen, CF3, C1-4 alkoxy, H0, N02, (un)substituted C1-4 alkyl, C1-4 alkylthio, (un)substituted C02H, etc.; R2

= Ph, naphthyl, heteroaryl, etc.; R3 = CN, CO2H, carboxylate ester, (un)substituted carboxamoyl, etc.; R4 = (un)substituted amino, NHCOR12, N(COR12)2, N:CHOCH2R12; R12 = H (un)substituted C1-4 alkyl, cyclic imido, Q; X = C2-4 alkylene, NHSO2R14; R14 = C1-4 alkyl, (un)substituted Ph; n = 0-2; R1 is located on ring positions 5-10], which demonstrate an antiproliferative effect on cell division and are useful in the treatment of diseases where excess cell proliferation or enzyme release is an aspect of the pathol. (no data), are prepared by the cyclization of R1-substituted 1-naphthols with NC(R3)C:CHR2. Thus, 1-naphthol was reacted with 3-(trifluoromethyl)benzylidenemalononitrile, producing I [R1 = H, R2 = 3-F3CC6H4, R3 = CN, R4 = NH2, n = 1].

MSTR 1

G2 = NH2

G4 = quinolinyl (opt. substd. by 1 or more G6)

G6 = 48 / alkylamino < containing 1-4 C>

C(O)-G2

Derivative: or salts Patent location: claim 1

Note: substitution is restricted

L5 ANSWER 125 OF 131 MARPAT COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 118:212755 MARPAT

TITLE: Preparation of cephalosporin compounds

INVENTOR(S): Takamura, Norio; Saito, Kunio; Matsushita, Tadahiro;

Yamaguchi, Totaro

PATENT ASSIGNEE(S): Tanabe Seiyaku Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 31 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 04261182	Α	19920917	JP 1991-287408	19910808
JP 06086461	В	19941102		
CA 2057129	A1	19930606	CA 1991-2057129	19911205

EP 544958 A1 19930609 EP 1991-311373 19911206 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE CN 1073444 A 19930623 CN 1991-111604 19911218 PRIORITY APPLN. INFO.: JP 1990-212040 19900809 GI

Ι

Cephalosporin compds. [I; R1 = NH2, etc.; R2 = OH, etc.; R2 = CO2H, etc.; R4 = H, alkyl, alkenyl, CH2R (wherein R = nucleophilic radical such as AcO, pyridino, quinolino, thiazolylthio, etc.); R5 = CO2H, etc.; R6 = H, etc.; dotted line = saturation or unsatn.], useful as broad-spectrum antibacterial agents, are prepared A solution of DMF and POC13 in CH2C12 was stirred at room temperature under Ar, cooled to -55° to -50°, treated with 13 g acid II (preparation given) in CH2C12 at -60° to -50°, and the solution was then treated with a suspension of MeC(OSiMe3):NSiMe3 and 5.43 g (syn)-I [R1 = Ph3CNH, R2 = 8-Ph2CHO, R3 = Ph2CHO2C, R4 = AcOCH2, R5 = CO2H, R6 = H, unsatd.]. The preferred dose was 5-40 mg/kg-day.

MSTR 1

G7 = 49

H₂C—G8

G8 = 80

G15 = CONH2 / NHCHO

Derivative:

and pharmacologically acceptable salts

Patent location: claim 1

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FILE 'REGISTRY' ENTERED AT 14:58:53 ON 27 AUG 2008

L1 STRUCTURE UPLOADED

L2 12 S L1 SAM

L3 208 S L1 FULL

FILE 'CA' ENTERED AT 14:59:23 ON 27 AUG 2008

L4 2 S L3

FILE 'MARPAT' ENTERED AT 14:59:47 ON 27 AUG 2008

L5 131 S L1 FULL

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10/572,914
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---Logging off of STN---

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STN INTERNATIONAL LOGOFF AT 15:11:02 ON 27 AUG 2008